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AKTIENGESELLSCHAFT; ().

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- (54) Title: ATP BINDING CASSETTE GENES AND PROTEINS FOR DIAGNOSIS AND TREATMENT OF LIPID DISORDERS AND INFLAMMATORY DISEASES
- (54) Titre: GENES ET PROTEINES DE CASSETTE DE LIAISON AVEC ATP, DESTINES AU DIAGNOSTIC ET AU TRAITEMENT DE DESORDRES LIPIDIQUES ET MALADIES INFLAMMATOIRES

(57) Abstract

Modulation of the activity of transmembrane proteins belonging to the ATP binding cassette (ABC) transporter protein family which are etiologically involved in cholesterol driven atherogenic processes and inflammatory diseases like psoriasis, lupus erythematodes and others provides therapeutic means to treat such diseases. Furthermore, detection of herein identified ABC transporter proteins of their respective biochemical activities involved in such atherogenic and inflammatory processes provides diagnostic means for clinical application of diagnosis and monitoring of dyslipidemias, atherosclerosis or inflammatory diseases like psoriasis and lupus erythematodes.

(57) Abrégé

Selon l'invention, la modulation de l'activité de protéines transmembranaires qui appartiennent à la famille de protéines de transport (ABC) de cassette de liaison avec ATP et sont impliquées de manière étiologique dans des processus athérogènes provoqués par le cholestérol et dans des maladies inflammatoires comme le psoriasis, le lupus érythémateux et autres, constitue un moyen thérapeutique de traiter de telles maladies. En outre, la détection des protéines de transport (ABC) ici identifiées et de leurs activités biochimiques respectives, impliquées dans de tels processus athérogènes et inflammatoires, constitue un moyen de diagnostic destiné à l'application clinique de diagnostic et de surveillance des dyslipidémies, de l'athérosclérose ou de maladies inflammatoires telles que le psoriasis ou le lupus érythémateux.



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Description

ATP binding cassette genes and proteins for diagnosis and treatment of lipid disorders and inflammatory diseases

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Background of the invention

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Reverse cholesterol transport mediated by HDL provides a "protective" mechanism for cell membrane integrity and foam cell formation and cellular cholesterol is taken up by circulating HDL or its precursor molecules. The precise mechanism of reverse cholesterol transport however is currently not fully understood and the mechanism of cellular cholesterol efflux and transfer from the cell surface to an acceptor-particle, such as HDL, is yet unclear. Certain candidate gene products have been postulated playing a role in the process of reverse cholesterol transport [1]. Apolipoproteins

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(e.g. ApoA-I, ApoA-IV), lipid transfer proteins (e.g. CETP, PLTP) and enzymes (e.g. LCAT, LPL, HL) are essential to exchange cholesterol and phospholipids in lipoprotein-lipoprotein and lipoprotein-cell interactions. Different plasma membrane receptors, such as SR-BI [2; 3]. HB1/2 [4], and GPI-linked proteins (e.g. 120 kDa and 80 kDa) [5] as well as the sphingolipid rich microdomains (Caveolae, Rafts) of

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the plasma membrane have been implicated being involved in the process of reverse cholesterol transport and the exchange of phospholipids. How these membrane-

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microdomains are organized is in the current focus of interest for the identification of therapeutic targets. In recent studies SR-BI function as receptor for uptake of HDL

into the liver and steroidogenic tissues could be demonstrated and the effectivity of this process is highly dependent on the phospholipid environment [2].

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Cholesterol and phospholipid homeostasis in monocytes/macrophages and other cells involved in the atherosclerotic process is a critical determinant in atherosclerotic vessel disease. The phagocytic function of macrophages in host defense, tissue remodelling, uptake and lysosomal degradation of atherogenic lipoproteins and membrane fragments or other lipid containing particles has to be balanced by effective release mechanisms to avoid foam cell formation. HDL mediated reverse

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cholesterol transport, supported by endogenous ApoE and CETP synthesis and secretion provides an effective mechanism to release excessive cholesterol from macrophages and other vascular cells.

Alternatively, reduced cholesterol and triglyceride/fatty acid absorption by intestinal mucosa cells as well as increased lipid secretion from hepatocytes into the bile will lower plasma lipids and the concentration of atherosclerotic lipoproteins.

Summary of the invention

New cholesterol responsive genes were identified with differential display method in human monocytes from peripheral blood that were subjected to macrophage differentiation and cholesterol loading with acetylated LDL and subsequent deloading with HDL₃.

In an initial screen ABCG1 (ABC8), a member of the rapidly growing family of ABC (ATP-Binding Cassette) transport systems, that couple the energy of ATP hydrolysis to the translocation of solutes across biological membranes, was identified as a cholesterol sensitive switch. ABCG1 is upregulated by M-CSF dependent phagocytic differentiation but expression is massively induced by cholesterol loading and almost completely set back to differentiation dependent levels by HDL₃.

In a more detailed analysis 37 already characterised ABC members and 8 Fragment - sequences (Table 2) were analysed in monocyte/macrophage cells by RT-PCR (linear range) for differentiation dependent changes and cholesterol sensitivity.

Among the 45 tested ABC-transporter genes 18 of the characterized ABC transporters and 2 of the Fragment -sequence based ABC-transporters are cholesterol sensitive (Example 4).

The cholesterol sensitive ABC-transporter are named according to the new ABC-

nomenclature and listed in Table 3 with the new and the old designations, respectively.

The most sensitive gene was ABCG1. ABCG1 is the human homologue of the drosophila white gene. Sequencing of the promoter of ABCG1 (Example 7) shows important transcription factor binding sites relevant for phagocytic differentiation and lipid sensitivity.

Antisense treatment of macrophages during cholesterol loading and HDL₃-mediated deloading clearly identified ABCG1 as a cholesterol transporter and the efflux of choline-containing phospholipids (phosphatidylcholine, sphingomyelin) was also modulated. Northern- and Western-blot analysis provided further support that inhibition of cholesterol transport is associated with lower ABCG1 mRNA expression and ABCG1 protein levels (Example 5).

Considerable evidence was derived from energy transfer experiments (Example 3) that ABCG1 in the cell membrane is in a regulated functional cooperation (e.g. cell differentiation, activation, cholesterol loading and deloading) with other membrane receptors that have either transport- (e.g. LRP-LDL receptor related protein) or signalling- and adhesion—function (e.g. integrins, integrin associated proteins) which is also supported by sequence homology of extracellular domains as well as other parts of the ABCG1 sequence. For example the protein sequence of the region of the third extracellular loop of ABCG1, i.e. aminoacid residues 580 through 644, shares homology with fibronectin (aa 317-327), integrinβ5 (aa 538-547), RAP (aa 119-127), LRP (aa 2874-2894), apoB-100 precursor (aa 4328-4369), glutathion-S-transferase (aa 54-78) and glucose transporter (aa 371-380). Sequence comparison of all cholesterol sensitive transporters indicates this as a general principle of ABC transporter function and regulation.

Among the other cholesterol sensitive gencs ABCA1 (ABC1) was further characterized. ABCA1 was identified in the mouse as an IL-1beta transporter

involved also in apoptotic cell processing. We show here, by RT-PCR (Table 2) and confirmation by Northern analysis, based on the newly detected human ABCA1 cDNA sequence (Example 6), that ABCA1 follows the same regulation as ABCG1.

Moreover, the ABCA1-knockout mice (ABCA1-/-) show massively reduced levels of serum lipids and lipoproteins. The expression of ABCA1 in mucosa cells of the small intestine and the altered lipoprotein metabolism in ABCA1-/- mice allows the conclusion that ABCA1 plays a major role in intestinal absorption and translocation of lipids into the lymph-system

Analysis of genetic defects that affect macrophage cholesterol homeostasis identified dysregulated ABCA1 as a gene locus involved in the HDL-deficiency syndrome (Tangier-Disease). This disease is associated with hypertriglyceridemia and splenomegaly.

Another as yet not described HDL-deficiency syndrome associated with early onset of coronary heart disease and psoriasis showed a dysregulation of the chromosome 17 associated ABC-sequences (ABCC4 (MRP3); ABCC3 (MRP3); ABCA5 (Fragment 90625); ABCA6 (Fragment 155051) :17q21-24). This points to an association with the predicted gene locus for psoriasis at chromosome 17.

A recently sequenced human ABC-transporter (ABCA8, Example 9) shows high homology to ABCA1 and also belongs to the group of cholesterol sensitive ABC-transporter.

ABCC5 (MRP5, sMRP) is a member of the MRP-subfamily among which ABCC2 (MRP2, cMOAT) was characterized as the hepatocyte canalicular membrane transporter that is involved in bilirubin glucoronide secretion [9] and identified as the gene locus for Dubin-Johnson Syndrome [10] a disorder associated with mild chronic conjugated hyperbilirubinemia.

Furthermore, the identification of ABCA1 as a transporter for IL-1 β identifies this gene as a candidate gene for treatment of inflammatory diseases including rheumatoid arthritis and septic shock. The cytokine IL-1 β is a broadly acting proinflammatory mediator that has been implicated in the pathogenesis of these diseases.

Moreover, we could demonstrate, that glyburide as an inhibitor of IL-1 β secretion inhibits not only Caspase I mediated processing of pro-IL-1 β and release of mature IL-1 β but simultaneously inhibits ceramide formation from sphingomyelin mediated by neutral sphingomyelinase and thereby releases human fibroblasts from G_2 -phase cell cycle arrest. These data provide a further mechanism indicative for a function of ABCA1 in signalling and cellular lipid metabolism.

Autoimmune disorders that are associated with the antiphospholipid syndrome (e.g. lupus erythematodes) can be related to dysregulation of B-cell and T-cell function, aberrant antigen processing, or aberrations in the asymmetric distribution of membrane phospholipids. ABC-transporters are, besides their transport function, candidate genes for phospholipid translocases, floppases and scramblases that regulate phospholipid asymmetry (outer leaflet: PC+SPM; inner leaflet: PS+PE) of biological membranes [11]. There is considerable evidence for a dysregulation of the analysed ABC-transporters in patient cells. We conclude that these ABC-cassettes are also candidate genes for a genetic basis of antiphospholipid syndromes such as in Lupus erythematodes.

In summary, the ABC genes ABCG1, ABCA1 and the other cholesterol-sensitive ABC genes as specified herein, can be used for diagnostic and therapeutic applications as well as for biochemical or cell-based assays to screen for pharmacologically active compounds which can be used for treatment of lipid disorders, atherosclerosis or other inflammatory diseases. Thus it is an objective of the present invention to provide assays to screen for pharmacologically active compounds which can be used for treatment of lipid disorders, atherosclerosis or

other inflammatory diseases. Further the invention provides tools to identify modulators of these genes and gene products. These modulators can be used for the treatment of lipid disorders, atherosclerosis or other inflammatory diseases or for the the preparation of medicaments for treatment of lipid disorders, atherosclerosis or other inflammatory diseases. The medicaments comprise besides the modulator acceptable and usefull pharmaceutical carriers.

Abbreviations

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		aa	Amino acid
		ABC	ATP-binding cassette
		ABCA#	ATP-binding cassette, sub-family A (ABC1), member #
15		ABCB#	ATP-binding cassette, sub-family B (MDR/TAP), member #
		ABCC#	ATP-binding cassette, sub-family C (CFTR/MRP), member #
		ABCD#	ATP-binding cassette, sub-family D (ALD), member #
20		ABCE#	ATP-binding cassette, sub-family E (OABP), member #
	•	ABCF#	ATP-binding cassette, sub-family F (GCN20). member #
		ABCG#	ATP-binding cassette, sub-family G (WHITE), member #
25		ABCR	Homo sapiens rim ABC transporter
25		AcLDL	Acetylated LDL
		ADP1	ATP-dependent permease
		ALDP	Adrenoleukodystrophy protein
30		ALDR	Adrenoleukodystrophy related protein
		ApoA	Apolipoprotein A
		ApoE	Apolipoprotein E
35		ARA	Anthracycline resistance associated protein
		AS	Antisense
		ATP	Adenosine triphosphate
		CETP	Cholesteryl ester transfer protein
40		CFTR	Cystic fibrosis transmembrane conductance regulator
		CGT	ceramide glucoxyl transferase
		CH	Cholesterol
45		cMOAT	Canalicular multispecific organic anion transporter
		dsRNA	Double stranded RNA
		Fragment	Gen Fragment
		FABP	plasma membrane fatty acid binding protein
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	FACS	Fluorescence activated cell sorter
	FATP	intracellular fatty acid binding protein
10	FCS	foetal calve serum
	FFA	free fatty acids
	GAPDH	Glyceraldehyde-3-phosphate dehydrogenase
	GCN20	protein kinase that phosphorylates the alpha-subunit of translation
15		initiation factor 2
	GPI	Glycosylphosphatidylinositol
	HaCaT	keratinocytic cell line
20	HDL	High density lipoprotein
	HL	Hepatic lipase
	HllyB	haemolysin translocator protein B
25	HMT1	yeast heavy metal tolerance protein
20	HPTLC	High performance thin layer chromatography
	IL	Interleukin
	LCAT	Lecithin:cholesterol acyltransferase
30	LDL	Low density lipoprotein
	LPL	Lipoprotein lipase
	LRP	LDL receptor related protein
35	MDR	Multidrug resistance
	MRP	Multidrug resistance-associated protein
	PC	Phosphatidylcholine
1	PE	Phosphatidylethanolamin
40	PL	Phospholipid
	PLTP	Phospholipid transferprotein
	PMP	peroxisomal membrane protein
45	PS	Phosphatidylserine
	RNA	Ribonucleic acid
	RT-PCR	Reverse transcription - polymerase chain reaction
50	SDS	Sodium dodecyl sulfate
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SL

Sphingolpid

sMRP

Small form of MRP

SPM

Sphingomyelin

SR-BI

Scavenger receptor BI

SUR

Sulfonylurea receptor

TAP

Antigen peptide transporter

TG

Triglycerides

TSAP

TNF-alpha stimulated ABC protein

UTR

untranslated region

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Description of the Figures

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Figures 1 to 5 are showing nucleotide and protein sequences described in this application. The sequences are repeated in the sequence listing.

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Description of Tabels:

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Table 1:

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Levels of RNA transcripts of ABCG1 (ABC8), ABCA1 (ABC1) and ABCA8 in human tissues were determined by Northern blot analysis of a multiple tissue dot-blot (Human RNA MasterBlot, Clontech Laboratories, Inc., CA, USA). The relative amount of expression is indicated by different numbers of filled circles.

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Table 2:

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The expression pattern of ABC-transporters in monocytes, monocyte derived macrophages (3 days cultivated monocytes in serum free Macrophage-SFM medium containing 50 ng/ml M-CSF), AcLDL incubated monocytes (3 days with 100 µg/ml) followed by HDL₃ (100 µg/ml) incubated monocytes is shown. Expressed genes are tested for cholesterol sensitivity by semiquantitative PCR.

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For known ABC-Transporter the chromosomal location and the transported molecules are also presented.

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Table 3:

Disorders, that are associated with ABC-transporters are shown. The chromosomal location is indicated and the relevant accession number in OMIN (Online Mendelian Inheritance in Man).

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Table 4:

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Expression of ABC-Transporters in HaCaT keratinocytic cells during differentiation

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Table 1

Tissue	ABCG1	ABCA1
	(ABC8)	(ABC1)
Adrenal gland	••••	•••
Thymus	••••	••
Lung	••••	•••
Heart	•••	••
Skeletal	••	•
Brain	•••	••
Spleen	••••	••
Lymphnode	•••	•
Pancreas	•	•
Placenta	••••	••••
Colon	. ••	•
Small intestine	••	••••
Prostate	••	•
Testis	• **	•
Ovary	••	•
Uterus	• • •	••
Mammary gland	••	•
Thyroid gland	••	••
Kidney	••	•
Liver	•••	•••
Bone marrow	•	•
Peripheral leukocytes	•	•
Fetal tissue		
Fetal brain	•	••
Fetal liver	-92 • 32	••••
Fetal spleen	••	•••
Fetal thymus	••	••
Fetal lung	••	•••

Table 2: Cholesterol dependent gene regulation of human ABC transporters

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Gene		chromosomal	peripheral blood	3 days old M-CSF	cholesterol loading	cholesterol deloading	transported
·		localization	monocytes	MCSF MCi	(acLDL)	(HDL3)	molecules
ABCGI	(ABC8)	21q22.3	+	1	11	- 11	cholesterol / choline PL
ABCA1	(ABCI)	9922-31	.+	1	↑ ↑.	11	cholesterol / IL-10
ABCC5	(MRP5)	3q25-27	+ 4	1	1 1.	Ţ	=
ABCD1	(ALDP, ALD)	Xq28	+	, 1	1	1	very long chain fatty acid
ABCA5	(est90625)	17q21-25	+	1	1	+	
ABCBII	(BSEP, SPGP)	2q24	+	1	↑ ↑	J .	bile acids
ABCA8	(ABC-new)		+	+	1	Ţ	
ABCC2	(MRP2)	10q23-24	+	+	1	1	bilirubin glucuronide
ABCB6	(est45597)	2q33-36	+	+	1	1	
ABCC1	(MRP1)	16p13.12	+ .	1	1	—	eicosanoids
ABCA3	(ABC3)	16p13.3	+	1	1	nr	
est113353	0	2.1	+ ·	1	1	nr	
ABCB4	(MDR3)	7q21	+	<u> </u>	↓	1	phosphatidylcholine
ABCG2 (st157481,ABCP)	4q22-23	+	1	1	T	
ABCC4	(MRP4)	13q31	+	1	1	T	
ABCB9	(est122234)	12q24	+	↑	1	1	
ABCD2	(ALDR)	12q11	+	1		1	very long chain fatty acid
ABCB1	(MDR1)	7q21	+	+	-	1	phospholipids,amphiphile
ABCA6	(est155051)	17q21	+	1	1	nr .	
est640918		100	+	1	1	nr	
ABCD4	(P70R)	14q24.3	+	1	nr	nr	
ABCA2	(ABC2)	9q34	+	↑	nr .	nr	
ABCF2	(est133090)	7q35-36	+	1	nr	nr	
ABCB7	(ABC7)	Xq13.1-3	+	1	nr	nr	iron
ABCF1	(ABC50,TSAP)	6p21.33	+	1	nr	nr	
ABCC6	(MRP6)	16p13.11	+	 +	nr	nr	*
ABCB5	(est422562)	7p14	+		nr	nr	
ABCC3	(MRP3)	17q11-21	+	nr	ពក	nr	
АВСЛ4	(ABCR)	1 p22	-1	nr	nr	nr	retinoids, lipofuscin
ABCB2	(TAP1)	6p21.3	+	nr	nr	nr	peptides
ABCB3	(TAP2)	6p21.3	+	nг	nr	nr	peptides

Gene		chromosomal localization	peripheral blood monocytes	3 days old M-CSF MD	cholesterol loading (acLDL)	cholesteroi deloading (HDL3)	transported molecules
ABCF3	(est201864)	3q25.1-2	+	nr	nr	nr	
ABCB8	(est328128)	7q35-36	+	T	nr	nr	
ABCE1	(OABP)	4q31	+	1	nr	nr	
ABCB10	(est20237)	1q32	+	1	nr	nr	
est698739			+	1,	nr	nr	
ABCC10	(est182763)	6p21	+	ηr	nr	nr	
ABCC7	(CFTR)	7q31	Ø.	Ø	Ø	Ø	ions
ABCC8	(SUR-1)	11p15.1	Ø	Ø	Ø	Ø	
ABCD3	(PMP70)	1p21-22	Ø	Ø	Ø	Ø	
Huwhite2			Ø	Ø	Ø	Ø	
est1125168			Ø	Ø	Ø	Ø	
est 1203215			Ø	Ø	Ø	Ø	
est I 68043			Ø	Ø	Ø	Ø	
est990006	<u>-</u>		Ø	Ø	Ø	Ø	· · · · · · · · · · · · · · · · · · ·

nr=not regulated

∩ = upregulated

U= downregulated

half (hs) or full size (fs) transporter as deduced from the mRNA size

Table 3

Disorders	Genomic	Associated gene	OMIM acc.nr
Metabolic disorders:	1	L.	acc.itt
Cystic fibrosis	7q31.3	ABCC7 (CFTR)	219700
Dubin Johnson syndrome (mild chronic conjugated hyperbilirubinemia)	10q24	ABCC2 (CMOAT)	237500
Progressive familial intrahepatic cholestasis type III (PIFC3)	7q21.1	ABCB4 (MDR3)	602347
Byler disease (PF1C2)	2q24	ABCB11 (BSEP, sPGP)	601847
Familial persistent hyperinsulinemic hypoglycemia	11p15.1	ABCC8 (SUR-1)	601820
IDDM	6p21.3	ABCB2 (TAP1)/ABCB3 (TAP2)	222100
Neuronal disorders:			
Adrenolcukodystrophy	12q11	ABCD2 (ALDR)	300100
Zellweger's syndrome	1p22-21	ABCD3 (PMP70)	214100
Multiple Sclerosis	6p21.3	ABCB2 (TAP1)/ABCB3 (TAP2)	126200
X-linked Sideroblastic anemia with spinoccrebellar ataxia	Xq13.1-3	ABCB7 (ABC7)	301310
Menkes disease (altered homeostasis of metals)	Xq13	ABCB7 (ABC7)	309400
Immune/Hemostats disorders:		L	
Herpes simplex virus infection [12]	6p21.3	ABCB2 (TAP1)/ABCB3 (TAP2)	
Behcet's syndrome	6p21.3	ABCB2 (TAP1)/ABCB3 (TAP2)	109650
Bare lymphocyte syndrome type I	6p21.3	ABCB2 (TAP1)/ABCB3 (TAP2)	209920
Scott syndrome	7q21.1	ABCBI (MDRI)	262890
Retinal dystrophies:			
Fundus flavi maculatus with macular dystrophy	lp13-21	ABCA4 (ABCR)	601691
uvenile Stargardt disease	1p13-21	ABCA4 (ABCR)	248200
Age-related macular degeneration	1p13-21	ABCA4 (ABCR)	153800
Cone-rod dystrophy	1p13-21	ABCA4 (ABCR)	600110
Retinitis pigmentosa	1p13-21	ABCA4 (ABCR)	601718

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Diseases with evidence for involvement of		Assumed gene	1
ATPcassettes/translocases and floppases[80]			
BRIC	[18]	Assumed	243300
(Benign recurrent intrahepatic obstructive jaundice)		,	
Psoriasis	17q11-12	ABCA5	602723
	17q21-24	(Fragment	177900
		90625)	601454
		ABCC3 (MRP3)	
Lupus erythematodes - Antiphospholipid Syndrome		Translocase	152700
	-	Flippase	
PFIC(Prog. Fatal familial intrahepatic choestasis) PFIC1	18q21-22	ATP	211600
		Transporters	
Neurological disorders mapped to gene locus of ABCG1 (Al	3C8)		L
Autosomal bipolar affective disorder	21q22.3	ABCG1 (ABC8)	125480
Autosomal recessive non-syndromic deafness	21q22.3	ABCGI (ABC8)	601072
Down Syndrome	21q22.3	ABCG1 (ABC8)	190685
(ABC-8 may be a candidate for the Brushfield spots -			
mottled, marble or speckled irides frequently seen in Down-			
Syndrome)			
Linkage to phosphofructokinase (liver type)	21q22		171860
HDL-deficiency syndromes,	9q31	ABCA1 (ABC1)	205400
Gen responsible for Tangier Disease			

Table 4: Expression of ABC-Transporters in HaCaT keratinocytic cells during differentiation

Gene	chrom, localisation	initial expression	differentiation dependent	known or putative
ABCG1 (ABCR)	21 q22.3	++++	· •	cholesterol choline-PL
	-	*		
ABCC3 (MRP3)	17 q11-q12	++++	↑	
ABCA8	19 P13	++++	↑	
ABCC1 (MRPI)	16 p13	++++	₹ ¥ (max. day 2)	PGA ₂ , LTC ₄
				DNP-SG
ABCD4 (PMP69, P70R)	14 q24	++++	7 ¥ (max. day 2,4)	<i>y</i> ×
ABCC2 (MRP2)	10 q24	†·!·+	7 № (max. day 2)	billrubin
		•		glucuronide
ABCA3 (ABC3)	16 p13	+	2 3 (max. day 4,6)	*
ABCA5 (ABCR)	1 p21	+	기 및 (max. day 4)	fetinoid,
	*	,		lipoluscin
ABCAI (ABCI)	9 q22-q31	+	オン (max. day 6)	
ABCC6 (MRP6)	16 p13.11	+	オ 3 (max. day 4)	-
ABCC4 (MRP4)	13 q31	++++	계 및 (max, day 2,4)	
ABCA2	9 q34	++++	7 3 (max. day 6)	
ABCC5 (MRP5, SMRP)	3 q27	+;;;;;	∌ (max. day 2,4)	

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ABCB6 (est45597)	T			
	2	++++	7 № (max. day 2,4)	
ABCB7 (ABC7)	X q13.1-3	1.1+++	# ≌ (max. day 4)	irons
TAPI (ABCBI)	6 p21.3	++++	기 및 (max. day 4,6)	peptides
TAP2 (ABCB2)	6 p21.3	++++	カン (max. day 2,4)	peptides
ABCB8 (cst328128)	7 q35-36.	+++++	게 및 (max day 2)	y 2
ESTG40918	17 q24	+	기 및 (max day 4)	-
ABCC7 (CFTR)	7 q31	+++	カン (max day 4)	
ABCB10 (est20237)	1 q32	+++	カン (max. day 2)	
ABCF1 (TSAP)	6 р21.33	++++	+	
ABCC10 (cst182763)	q32	++++	1	
ABCET (OABP)	4 q31	++++	J	* .
EST698739	17 q24	++++	•	
ABCF2 (est133090)	7 q35-q36	++++	ψ	
ALD (ABCDI,ALDP)	X q28	****	ψ	VLCFA
ABCA5 (est90625)	17 q21-q24	+++	ψ	-
ABCB5 (est422562)	7 pl4	++++	+	
ABCB9 (est122234)	12 424-q _{ter}	++	4	
ABCD2 (ALDR)	12 q11	+	Ψ	VLCFA
ABCF3 (cst201864)	3 q25.1-2	++++	4	
ABCG2 (ABC15,ABCP)	4 q22-q23	++++	ų i	-
EST1133530	4 pl6pter	++++	7 4	

Huwhite	11 q23	++++	ψ	
ABCA6 (cst155051)	17 q21	++	ψ	
BSEP (ABCBII,sPGP)	2 q24	+	↓ ↑ (max day 6)	
ABCB4 (MDR3)	7 q21	not expressed		phosphatidyl-
		*		choline
ABCD3 (PMP70)	1 p22	not expressed		
ABCBI (MDR1)	7 q21	not expressed		phospholipids amphiphiles
EST168043	2 p15-16	not expressed	0	* -
EST990006	17 q24	not expressed		
ABCC8(SURI)	t1 pt5.1	not expressed		

+: relative expression n.d.: not determined

↑: upregulated ↓: downregulated ≯ \(\mathbf{1} \): biphasic expression

Description of specific embodiments

Candidate gene identification during cholesterol loading and deloading of human monocyte derived macrophages

In order to discover genes that are involved in the cholesterol loading and/or deloading in vitro assays were set up. Particularly, gene expression in human blood derived monocytes and macrophages elicited by cholesterol and its physiological transport formulation, i.e. various low density lipoprotein (LDL) particle species like AcLDL, was studied.

Elutriated human monocytes were cultivated in M-CSF containing but serum free macrophage medium supplemented with AcLDL (100 μg protein/ml medium) for three days, followed by cholesterol depletion replacing AcLDL by HDL₃ (100 μg protein/ml medium) for twelve hours. Differential display screening for new candidate genes, regulated by cholesterol loading/deloading, was performed (Example 1).

Identification of a new cholesterol sensitive gene

ABCG1 (ABC8) was discoverd as a novel cholesterol sensitive genc. ABCG1 belongs to the ATP binding cassette (ABC) transporter gene family. ABCG1 was recently published as the human analogue of the drosophila white gene [6-8].

The gene is strongly upregulated by Acl.DL-mediated cholesterol loading, and almost completely downregulated by HDL₃ mediated-cholesterol deloading, as confirmed by Northern blot (Example 2). Nothern blot analysis oh mRNA from human monocyte-derived macrophages obtained from the peripherical blood probands clearly show upregulation of ABCG1 mRNA formation upon Acl.DL incubation. In sharp contrast, ABCG1 mRNA expression was decreased in such macrophages upon incubation with HDL₃ containing medium.

ABCG1 expression in cholesterol loaded and deloaded cells after four days predifferentiation

For effective cholesterol loading monocytes must be differentiated to phagocytic-macrophage like cells. During this period scavenger receptors are upregulated and promote AcLDL uptake leading to cholesteryl ester accumulation. After four days preincubation period we have incubated the cells for one, two and three days with AcLDL (100 µg/ml) to show cholesteryl ester accumulation. After two days of loading we deloaded the cells with HDL3 for 12 hours. 24 hours and 48 hours, respectively. ABCG1 is time dependently upregulated during the AcLDL loading period and downregulated by HDL3 deloading (Examples 2 and 3) In order to confirm time dependent increase of ABCG1 mRNA expression after AcLDL challenge in human monocyte derived macrophages, Nothern blot analyses for ABCG1 mRNA quantification were made, RNA samples from the macrophages were harvested at day zero and day four as controls and mRNA samples were taken one, two, and three days after AcLDL treatment of macrophages, which started at day four. A dramatic increase of ABCG1 mRNA content of the macrophages could be detected from day five through day seven by Nothern blot analyses.

This regulation shows the same pattern as changes of cellular cholesteryl ester content (Example3). Cholesterol ester accumulation starts in monocyte-derived macrophages upon AcLDL stimulation from a base level below 5 nmol/mg cell protein at day four up to 120 nmol/mg cell protein at day seven (i.e. three days after AcLDL application).

Tissue expression

Besides cholesterol loaded macrophages ABCG1 is prominently expressed in brain, spleen, lung, placenta, adrenal gland, thymus and fctal tissues (Table 1).

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Chromosomal location and associated genes and diseases

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The ABCG1 gene maps to human chromosome 21q 22.3. Also localized in this region 21q 22.3 are the following genes: integrin β 2 (CD18), brain specific polypeptide 19, down syndrome cell adhesion molecule, dsRNA specific adenosine deaminase, cystathionine β synthase, collagen VI alpha-2, collagen XVIII alpha-1, autosomal recessive deafness, and amyloid beta precursor.

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This chromosomal region is in close proximity to other regions involved in Down syndrome, autosomal dominant bipolar affective disorder, and autosomal recessive non-syndromic deafness.

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Extracellular loop of ABCG1 (ABC8) for antibody generation

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The putative structure of the hydrophobic transmembrane region of ABCG1 shows 6 transmembrane spanning domains, and 3 extracellular loops, two of them are 9- and 8-amino acids-long, respectively, while the third one is 66-amino acids-long.

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The larger one of the two intracellular loops consists of 30 amino acids. Similarity-survey in protein databases for homologies the 3rd extracellular loop (IIIex) with other genes resulted in the identification of fibronectin, integrin \$\beta\$5, RAP, LRP (LDL receptor related protein) apo-lipoprotein B 100 precursor protein, glutathion Stransferase and glucose transporter.

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A polyclonal antiserum was generated against the 3rd extracellular loop (IIIex) of ABCG1 in order to perform flow cytometric analysis, energy transfer experiments and Western-blotting (see Example 3). In the amino acid sequence of ABCG1 the 3rd extracellular loop (IIIex) comprises 66 amino acids comprises 66 amino acids from amino acid 580 through 644. The peptide fragment for antibody generation comprises the amino acid residues 613 through 628 of ABCG1 polypeptide. ABCG1 obviously interacts with endogenous sequence motivs with other membrane receptors

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involved in transport (e.g. LRP, RAP), signalling and adhesion (e.g. integrins, integrin associated proteins) as a basis of ABCG1-function and regulation. Moreover sequence comparisons of all ABC-transporters listed in Table 3 indicates functional cooperation with other membrane receptors as a general principle of the whole gene family.

Subfamily-Analysis

Evolutionary relationship studies with the whole ABC transporter family have shown that ABCG1 (ABC8) forms a subfamily together ABCG2 (cst157481) and this subfamily is closely related to the full-size transporters ABCA1 (ABC1). ABCA2 (ABC2), ABCA3 (ΛBC3), ABCA4 (ΛBCR) and the half-size transporter ABCF1 (TSAP).

Recent studies by Allikmets et al. have identified 21 new genes as ABC transporters by expressed sequence tags database search [13].

General description of the ABC transporter family

The ATP-binding cassette (ABC) transporter superfamily contains some of the most functionally diverse proteins known. Most of the members of the ABC family (also called traffic ATP-ases) function as ATP-dependent active transporters (Table 3). The typical functional unit consists of a pair of ATP-binding domains and a set of transmembrane (TM) domains. The TM-domains determine the specificity for the type of molecule transported, and the ATP-binding domains provide the energy to move the molecule through the membrane [14; 15]. The variety of substrates handled by different ABC-transporters is enormous and ranges from ions to peptides. Specific transporters are found for nutrients, endogenous toxins, xenobiotics, peptides, aminoacids, sugars, organic/inorganic ions, vitamins, steroid hormones and drugs [16; 17].

ABC-transporter associated diseases

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The search for human disease genes (Table 3) provided a number of previously undiscovered ABC proteins [16]. The best characterized disease caused by a mutation in an ABC transporter is cystic fibrosis (ABCC7 (CFTR)). Inherited disorders of peroxisomal metabolism as Adrenoleukodystrophy and Zellweger's syndrome also show alterations in ABC transporters. They are involved in peroxisomal beta-oxidation, necessary for very long chain fatty acid metabolism [18].

Antisense against ABCGI inhibits cholesterol efflux to HDL,

Since ABCG1 is a cholesterol sensitive gene and other ABC transporters are known to be involved in certain lipid transport processes, the question arises whether ABCG1 plays a role in transport of cholesterol, phospholipids, fatty acids or glycerols. Therefore antisense experiments were performed to test the influence of ABCG1 on lipid loading and deloading. The inhibition of ABCG1 with specific antisense oligonucleotides decreased the efflux of cholesterol and phosphatidyl-choline to HDL₃. (Example 5)

20 Other cholesterol sensitive ABC transporter

Cloning and sequencing of the human ABCA1 (ABC1) provided the information to characterize ABCA1 for cholesterol sensitivity, and tissue distribution (Example 6). Another cholesterol sensitive human ABC transporter (ABCA8) has been cloned and sequenced (Example 8)

Characterization of the ABCG1 promoter region

The ABCG1 promoter has the characteristic binding sites for transcription factors that are involved in the differentiation of monocytes into phagocytic macrophages. The cholesterol sensitivity of the expression of ABCG1 is represented by the transcription factor pattern that is relevant for phagocytic differentiation (Example 7).

Examples

Example 1

Identification of cholesterol loading and deloading candidate genes

Monocyte isolation and cell culture

Monocytes were obtained from peripheral blood of healthy normolipidemic volunteers by leukapheresis and purified by counterflow clutriation. Purity of isolated monocytes was >95% as revealed by FACS analysis. 10×10^6 monocytes were seeded into 100 mm² diameters cell culture dishes under serum free conditions in macrophage medium for 12 hours in a humidified 37°C incubator maintained with a 5% CO2, 95% air atmosphere. After 12 hours medium containing unattached cells was replaced by fresh macrophage medium supplemented with 50 ng/ml human recombinant M-CSF (this medium is the standard medium for any further incubations).

Isolation of lipoproteins and preparation of AcLDL

Lipoproteins were prepared from human plasma from healthy volunteer donors by standard sequential ultracentrifugation methods in a Beckman L-70 ultracentrifuge equipped with a 70 Ti rotor at 4°C to obtain LDL (d=1,006 to 1,063 g/ml) and HDL, (d=1,125 to 1,21 g/ml). All densities were adjusted with solid KBr. Lipoprotein fractions are extensively dialyzed with phosphate-buffered saline (PBS) containing 5 mM EDTA. The final dialysis step was in 0,15 mol/L NaCl in the absence of EDTA. Lipoproteins were made sterile by filtration through a 0.45 μ m (pore-size) sterile filter (Sartorius).

LDL was acetylated by repeated addition of acetic anhydride followed by dialysis against PBS [19]. Modified LDL showed enhanced mobility on agarose gel electrophoresis.

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Incubation of monocyte-macrophages with AcLDL and HDL,

After 12 hours of preincubation cells were grown in the presence or absence (control) of 100 µg protein /ml AcLDL for further 3 day in medium. Then, the incubation medium was replaced with fresh medium and incubated with or without the addition of HDL₃ (100 µg/ml) for another 12 hours.

Differential display

Differential display screening was performed for new candidate genes that are regulated by cholesterol loading/deloading as described [20; 21]. In brief, 0,2 µg of total RNA isolated from monocytes at various incubations was reverse transcribed with specific anchored oligo-dT primers, using a commercially available kit (GeneAmp RNA PCR Core Kit, Perkin Elmer, Germany). The oligo-dT primers used had two additional nucleotides at their 3' end consisting of an invariable A at the second last position (3'-end) and A, C, G or T at the last position to allow a subset of mRNAs to be reverse transcribed. Here, a 13-mer oligo-dT (T101: 5'T11AG-2') was used in a 20-µl reaction at 2,5 µM concentration. One tenth of the cDNA was amplified in a 20-µl PCR reaction using the same oligo-dT and an arbitrary 10-mer upstream primer (D20 5'-GATCAATCGC-3'), 2,5 µM each, using 2,5 units of TAQ DNA Polymerase and 1.25 mM MgCl2. Amplification was for 40 cycles with denaturation at 94°C for 30 sec, annealing at 41°C for 1 min and elongation at 72°C for 30 sec with a 5 min extension at 72°C following the last cycle. All PCR reactions were carried out in a Perkin Elmer 9600 thermocycler (Perkin Elmer, Germany). PCR-products were separated on ready to use 10% polyacrylamide gels with a 5% stacking gel (CleanGel Large-10/40 ETC, Germany) under non-denaturating conditions using the Multiphor II electrophoresis apparatus (Pharmacia, Germany). The DNA fragments were visualized by silverstaining of the gel as previously described [22].

Cloning and sequencing of differentially expressed cDNAs

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cDNA bands of interest were cut out of the gel and DNA was isolated by boiling the gel slice for 10 min in 20 µl of water. A 4 µl aliquot was used for the following PCRreaction in a 20µl volume. The cDNA was reamplified using the same primer set and PCR conditions as above, except, that the final dNTP concentration was 1mM each. Reamplified cDNAs were cloned in the pUC18-vector using ABCC8 (SUR)eClone-Kit (Pharmacia), sequenced on an automated fluorescence DNA sequencer using the AutoRead Sequencing Kit (Pharmacia, Germany) and used as probes for Northern blot analysis [23].

Example 2

Northern Blot analyses of monocytes and macrophages after 3 days AcLDL incubation followed by 12 hours HDL3 incubation

Elutriated monocytes were incubated with AcLDL (100 µg/ml medium) for 2.5 days or differentiated for the same time without the addition of AcLDL as control. ABCG1 (ABC8) expression is 4 times stronger upregulated with AcLDL incubation than in differentiated monocytes .After the AcLDL incubation period cells were washed and incubated with HDL, for the next 12 hours or with medium alone as control. ABCG1 expression is almost completely downregulated by HDL3 incubation and only moderatly decreased in control incubation as confirmed by Northern blot. For effective cholesterol loading monocytes must be differentiated to macrophage like cells. During this period scavenger receptors are upregulated and promote AcLDL uptake leading to cholesteryl ester accumulation. To differentiated the cells prior to AcLDL-dependent cholesterol loading, we cultured the cells for four days in standard medium. At day four, cells were washed and incubated with AcLDL (100µg/ml medium) or in the absence of AcLDL as control for further one, two and three days to load the cells with cholesterol. At each timepoint cells were lysed with 0.1 % SDS and lipid was extracted as described in materials and methods and cellular cholesteryl ester was determined by HPTLC-separation. Cells were loaded time

dependently up to 120 nmol/mg cell protein after 3 days AcLDL loading, whereas in unloaded cells no choicsteryl ester accumulation could be observed.

To distinguish HDL₃ dependent and independent cholesterol efflux cells were pulsed with AcLDL (100 μg/ml) for three days with the coincubation of ¹⁴C-cholesterol (1,5 μCi/ml medium). Cells were washed and deloaded with HDL₃ (100 μg/ml) for 12 hours, 24 hours and 48 hours, respectively. Cells were incubated without the addition of exogenous lipid-acceptors as a control. After chase period the content of ¹⁴C-cholesterol was determined in the medium and in the cells by liquid scintillation as described in material and methods. The efflux of cholesterol is expressed in percent of cellular DPMs of total DPMs (counts in the cells plus medium) With HDL₃ the efflux is faster and more intense, than the efflux without the addition of HDL₃ as an endogenous lipid acceptor. After 12 hours cellular cholesterol content was reduced to 68 % with HDL₃-dependent deloading, and 86 % in HDL₃-independent deloading. After 48 hours only 35 % of loaded 14C-cholesterol was observed in the cells treated with HDL₃. In contrast, 70 % of loaded ¹⁴C-cholesterol was found in untreated cells

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In AcLDL pulsed cells the RNA-expression of ABCG1 is upregulated whereas no upregulation appears in the cells that were not loaded with AcLDL. Cells that were loaded for two days with AcLDL were deloaded with HDL₃ for 12, 24 and 48 hours (12h; 24h; 48h), and in the absence of exogenous lipid acceptors. The RNA-expression is downregulated again, in HDL₃ treated cells more intense than in cells treatet without any exogenous lipid acceptor.

Materials:

Macrophage medium (Macrophage-SFM) was obtained from Gibco Life Technologies, Germany. Human recombinant M-CSF was obtained from Genzyme Diagnostics, Germany, and antisense phosphorothioate oligonucleotides were supplied by Biognostics, Germany. All other chemicals were purchased from Sigma. Nylon membranes and a32P-dCTP were obtained from Amersham, Germany, 14C-

cholesterol and 3H-choline chloride from NEN, Germany, and cell culture dishes are Becton Dickinson, Germany

Isolation of total RNA and northern blotting

Total RNA was isolated at each time-point, before and after AcLDL incubation, and after HDL₃ incubation, respectivly, Washed cells were solubilized in guanidine isothiocyanate followed by sedimentation of the extract through cesium chloride [24]. For Northern analysis, 10 µg/lane of total RNA samples were fractionated by electrophoresis in 1,2% agarose agarose gel containing 6% formaldehyde and blotted onto nylon membranes (Schleicher & Schüll, Germany). After crosslinking with UV-irradiation (Stratalinker model 1800, Stratagene, USA), the membranes were hybridized with a cDNA probe for ABCG1 (ABC8). Hybridization and washing conditions were performed as recommended by the manufacturer of the membrane.

Example 3

Westernblot analysis of monocytes and macrophages after cholesterol loading and deloading

Protein expression of ABCG1 (ABC8) is upregulated in AcLDL-loaded and down-regulated in IIDL₃-deloaded monocyte-derived macrophages. Western blotting with a peptide antibody against ABCG1 as described in materials and methods is performed with 40 µg of total protein for each lane of SDS-PAGE. ABCG1-protein expression is shown in freshly isolated monocytes (day zero) and in differentiated monocytes (day four). From day four to day seven (5d; 6d; 7d) monocyte-derived macrophages were loaded with AcLDL or without AcLDL as control. AcLDL loaded cells from day 6 (6d) were deloaded with HDL₃ for 12, 24, and 48 hours and without exogenous added HDL lipid-acceptor. AcLDL increases the protein-expression, whereas HDL₃ decreases the expression to normal levels again.

Protein isolation and determination

At each timepoint cells were lysed with 0.1% SDS and the protein content was determined by the method of Lowry et al. [25].

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Generation of ABCG1 specific antibodies

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ABCG1 specific peptide antibodies were generated by immunization of chickens and rabbits with a synthetic peptide (Fa. Pineda, Berlin). The peptide sequence was chosen from the extracellular domain exIII amino acid residues 613-628 of ABCG1 comprising the amino acids REDLHCDIDETCHFQ (see sequence listing ID No. 53). After 58 days of immunization western blotting was performed with 1:1000

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diluted scrum and 1:10000 secondary peroxidase labelled antibody.

Electrophoresis and immunoblotting

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SDS-polyacrylamide gelelectrophoresis was performed with $40\mu g$ total cellular 15 protein per lane. Proteins were transferred to Immobilon as reported. Transfer was

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confirmed by Coomassie Blue staining of the gel after the electroblot. After blocking for at least 2 hours in 5% nonfat dry milk the blot was washed 3 times for 15 minutes in PBS. Antiserum generated as described was used at 1:1000 dilution in 5% nonfat

dry milk in PBS. The blot was incubated for 1 hour. After 4 times washing with PBS at room-temperature a secondary peroxidase-labelled rabbit anti chicken IgG-

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antibody (1:10000 diluted, Sigma) was incubated in 5% nonfat dry milk in PBS for 1 hour. After 2 times washing with PBS, detection of the immune complexes was

carried out with the ECL Western blot detection system (Amersham International

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PLC, UK).

Fluorescence resonance energy transfer:

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Monocytes were labelled with the specific antibodies for 15 minutes on icc, one antibody is labelled by biotin, the other one is labelled by phycocrythrin. After washing the cells were incubated with a Cy5-conjugated streptavidin for another 15 minutes.

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Distances between antibody labelled proteins on the cell surface is measured by energy transfer with a FACScan (Becton Dickinson). Following single laser excitation at 488 nm the Cy5 specific emmission represents an indirect excitation of Cy5 dependent on the proximity of the PE-conjugated antibody. The relative transfer efficiency was calculated following standardisation for the intensity of PE and Cy5 labelling and nonspecific overlap of fluorescence based on dual laser excitation and comparison to separately stained control samples.

Example 4

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Cholesterol sensitivity of ABCG1 (ABC8) and other members of the ABC-transporter family

The influence of cholesterol loading and deloading on other members of the ABC-family was also investigated to find out the potential second half-size ABC transporter.

Further analysis has been performed to examine the expression pattern of all human ABC transporters in monocytes and monocyte derived macrophages as well as in cholesterol loaden and deloaden mononuclear phagocytes.

The experiments were performed by RT-PCR with cycle-variation to compare the expression in the quantitative part of the distinct PCR. Primer sets were generated from the published sequences of the ABC-transporters. A RT-PCR with GAPDH primers was used as control.

Several ABC-transporters are also cholesterol sensitive which further supports the function of ABC-transporters in cellular lipid trafficking (Table 2).

Semi-quantitative RT-PCR

All known ABC-transporters are tested for AcLDL/HDL, sensitive regulation of expression using RT-PCR with cycle-variation to compare the expression in the

quantitative part of the distinct PCR. 1 µg of total RNA was used in a 40 µl reverse transcription reaction, using the Reverse Transkription System (Promega, Corp. WI, USA). Aliquots of 5 µl of this RT-reaction was used in 50µl PCR reaction. After denaturing for 1,5 min at 94°C, 35 or less cycles of PCR were performed with 92,3°C for 44s, 60,8°C for 40s (standard annealing temperature differs in certain primer-combinations), 71,5°C for 46s followed by a final 5-min extension at 72°C. The Primer sets were generated from the published sequences of the ABC-transporters. A RT-PCR with primers specific for GAPDH was performed as control.

The expression pattern of ABC-transporters in monocytes, monocyte derived macrophages (3 days cultivated monocytes in serum free macrophage-SFM medium containing 50 ng/ml M-CSF), AcLDL incubated monocytes (3 days with 100 μ g/ml) followed by HDL₃ (100 μ g/ml) incubated monocytes is shown in Table 2. Expressed genes are tested for cholesterol sensitivity by semi-quantitative PCR.

15 Example 5:

Functional analyses of the cholesterol sensitive ABCG1 (ABC8) transporter gene by antisense oligonucleotide experiments

Antisense experiments were conducted in order to address the question, that beyond being regulated by cholesterol loading and deloading ABCG1 is directly involved in lipid loading and deloading processes.

In various experiments antisense oligonucleotides decreased the efflux of cholesterol and phosphatidylcholine to HDL₃. During the loading period with AcLDL the cells were coincubated with 17 different antisense oligonucleotides. To measure the efflux of cholesterol and phospholipids the cells were pulsed in the loading period with 1,5 μCi/ml ¹⁴C-cholesterol and 3μCi/ml ³H-choline chloride. The medium was changed and during the chase period cells were incubated with or without HDL₃ for 12 hours. The ¹⁴C-cholesterol and ³H-choline content in the medium and in the cell lysate was measured and the efflux was determined in percent of total ¹⁴C-cholesterol and ³H-choline loading.

The most effective antisense oligonucleotide (AS Nr.2) inhibited cholesterol and

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phospholipids efflux relative to cells that were treated with control antisense (AS control). A dose dependent decrease in cholesterol efflux of 16,79% (5nmol AS) and 32,01% (10 nmol AS) could be shown, respectively.

Antisense incubation

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To inhibit the induction of ABCG1 cells were treated with three different antisense oligonucleotides targeting ABCG1 or one scrambled control-antisense oligonucleotide during the AcLDL-incubation period.

Determination of cholesterol and phosphatidylcholine efflux from monocytes in dependency of antisense oligonucleotide treatment

To measure the efflux of cholesterol and phospholipids the cells were pulsed in addition to AcLDL-incubation with 1,5 μCi/ml ¹⁴C-cholesterol and 3μCi/ml ³H-choline chloride. The medium was changed and in chase period the cells were incubated with or without HDL₃ for 12 hours. Lipid extraction was performed according to the method of Bligh and Dyer [26]. The ¹⁴C-cholesterol and ³H-choline content in the medium and in the cell lysate was measured by liquid scintillation counting and the efflux was determined in percent of total ¹⁴C-cholesterol and ³H-choline loading as described [27]

Computer analyses

DNA and protein sequence analyses were conducted using programs provided by HUSAR, Heidelberg, Germany: http://genius.embnet.dkfz-heidelberg.de:8080.

Example 6

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Complete cDNA sequence of the human ATP binding cassette transporter 1 (ABCA1 (ABC1)) and assessing the cholesterol sensitive regulation of ABCA1 mRNA expression

cDNA Cloning and Primary Protein Structure

We have cloned a 6880-bp cDNA containing the complete coding region of the human ABCA1 gene (Figure 8) The open reading frame of 6603 bp encodes a 2201-amino acid protein with a predicted molecular weight of 220 kDa. This protein displays a 94% identity on the amino acid level in an alignment with mouse ABCA1 and can therefore be considered as the human ortholog.

Tissue Distribution of ABCA1 mRNA Expression

In order to examine the tissue-specific expression of ABCA1 a multiple tissue RNA master blot containing poly A* RNA from 50 human tissues was carried out. Northern Blot analysis demonstrates the presence of a ABCA1 specific signal in all tissues. It is mostly prominent in adrenal gland, liver, lung, placenta and all fetal tissues examined so far (Table 1). The weakest signals are found in kidney, pancreas, pituitary gland, mammary gland and bone marrow.

Sterol Regulation of ABCA1 mRNA Expression

In order to determine the regulation of ABCA1 in monocytes/macrophages during cholesterol loading/depletion Northern Blot analysis was performed. The cloned 1000-bp DNA fragment derived from PCR amplification of RNA from five day differentiated monocytes with primers ABCA1 3622f (CGTCAGCACTCTGATGATGGCCTG-3') and ABCA1 4620r (TCTCTGCTATCTCCAACCTCA-3') was hybridized to Northern Blots containing RNA of differentially cultivated monocytes (figure 12) As can be seen in lanes one to five, the ABCA1 mRNA is increased during in vitro differentiation of freshly isolated monocytes until day five. Longer cultivation results in a total loss of

expression. When the cells were incubated in the presence of AcLDL to induce sterol loading (lanes 6-8) beginning at day four, a much stronger accumulation of mRNA can be detected in comparison to control cells (lanes 2-5). When these cells were cultured with HDL₃ as cholesterol acceptor for 12h, 24h and 48h (lanes 9-11) the ABCA1 signal significantly decreases with respect to control cells incubated in the absence of HDL₃ (lanes 12-14). Taken together, these results indicate that ABCA1 is a sterol-sensitive gene which is induced by cholesterol loading and downregulated by cholesterol depletion.

Cell culture.

Peripheral blood monocytes were isolated by leukapheresis and counterflow elutriation (19JBC). To obtain fractions containing >90% CD 14 positive mononuclear phagocytes, cells were pooled and cultured on plastic Petri dishes in macrophage SFM medium (Gibco BRL) containing 25 U/ml recombinant human M-CSF (Genzyme) for various times in 5% CO₂ in air at 37°C. The cells were incubated in the absence (differentiation control) or presence of AcLDL (100 μg/ml) to induce sterol loading. Following this incubation the cells were cultured in fresh medium supplemented with or without HDL₃ (100 μg/ml) for additional times in order to achieve cholesterol efflux from the cells to its acceptor HDL₃.

Preparation of RNA and Northern blot analysis.

Total cellular RNA was isolated from the cells by guanidium isothiocyanate lysis and CsCl centrifugation (Chirgwin). The RNA isolated was quantitated spectrophotometrically and 15 μg samples were separated on a 1.2% agarose-formaldehyde gel and transferred to a nylon membrane (Schleicher & Schüll). After crosslinking with UV-irradiation (Stratalinker model 1800, Stratagenc), the membranes were hybridized with a 1000 bp DNA fragment derived from PCR amplification with primers ABCA1 3622f and ABCA1 4620r, stripped and subsequently hybridized with a human β -actin probe. In order to determine the tissue-specific expression of ABCA1 a multiple tissue RNA master blot containing

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poly A* RNA from 50 human tissues was purchased from Clontech. The probes were radiolabeled with $[\gamma^{-32}P]dCTP$ (Amersham) using the Oligolabeling kit from Pharmacia. Hybridization and washing conditions were performed following the method described previously (Virca).

cDNA cloning of human ABCA1

Based on sequence information of mouse ABCA1 cDNA we designed primers for RT-PCR analysis in order to amplify the human ABCA1 (ABC1) cDNA. Approximately 1 μ g of RNA from five day differentiated mononuclear phagocytes was reverse transcribed in a 20 μ l reaction using the RNA PCR Core Kit from Perkin Elmer. An aliquot of the cDNA was used in a 100 μ l PCR reaction performed with Amplitaq Gold (Perkin Elmer) and the following primer combinations: (primer names indicate the position in the corresponding mouse cDNA sequence):

mABC1-144f (5'-CAAACATGTCAGCTGTTACTGGA-3') and mABC1-643r (5'-TAGCCTTGCAAA-AATACCTTCTG-3'),

15 mABCI-1221f (5'-GTTGGAAAGATTCTCTATACACCTG-3') and mABCI-1910r (5'-CGTCAGCACTCTGATGATGGCCTG-3'), mABCI-3622f (5'-TCTCTGCTATCTCCAACCTCA-3') and mABCI-4620r (5'-ACGTCTTCACCAGGTAATCTGAA-3'), mABCI-5056f (5'-CTATCTGTGTCATCTTTGCGATG-3') and

mABC1-5857r (5'-CGCTTCCTCCTATAGATCTTGGT-3'),
mABC1-6093f (5'-AAGAGAGCATGTGGA-GTTCTTTG-3') and
mABC1-7051r (5'-CCCTGTAATGGAATTGTGTTCTC-3'),
hABC1-540f (5'-AACCTTCTCTGGGT1'CCTGTATC-3') and
hABC1-1300r (5'-AGTTCCTGGAA-GGTCTTGTTCAC-3'),
hABC1-1831f (5'-GCTGACCCCTTTGAGGACATGCG-3') and

hABC1-3701r (5'-ATAGGTCAGCTCATGCCCTATGT-3'),
hABC1-4532f (5'-GCTGCC-TCCTCCACAAAGAAAAC-3') and
hABC1-5134r (5'-GCTTTGCTGACCCGCTCC-TGGATC-3'),
hABC1-5800f (5'-GAGGCCAGAATGACATCTTAGAA-3') and
hABC1-6259r (5'-CTTGACAACACTTAGGGCCACAAT-3').

All PCR products were cloned into the pUC18 plasmid vector and the nucleotide sequences were determined on a Pharmacia ALF express sequencer using the dideoxy chain-termination method and fluorescent dye-labeled primers.

Example 7

Identification of the 5'end of ABCG1

We could partially prove the 5'-end of ABCG1 published by Chen [7] that differs from the 5'-end published by Croop [6] obtained from the mRNA of human monocytes/macrophages using a 5' RACE approach. In detail the sequence according to Chen et al. downstream of position 25 was in agreement with our own data. In contrast, our identified sequence differs from the one reported by Chen [7] and Croop [6] at a site upstream of position 25 (Chen [7]). The sequence SEQ ID NO: 32 shows the newly identified 5'-end followed by the sequence published by Chen [7] from position 25.

Molecular cloning and characterisation of the ABCG1 5'UTR

We identified several fragments by screening of a λ phage library which contained a total of app. 3 kb of the 5' UTR upstream sequence of the human ABCG1 gene. The

sequence that comprises the 5'UTR and part of exon 1 (described above) are given in SEQ ID NO: 54.

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The promoter activity of this sequence was proven by luciferase reporter gene assays in transiently transfected CHO cells.

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Putative transcription factor binding sites within the promoter region with the highest likelihood ratio for the matched sequence as deduced from the TransFac database, GFB, Braunschweig, Germany. Multiple binding sites for SP-1, AP-1, AP-2 and CCAAT-binding factor (C/EBP family) are present within the first 1 kb of the putative promoter region.

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Additionally, a transcription factor binding site involved in the regulation of apolipoprotein B was identified.

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Example 8

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Characterization of the human ABCA8 full length cDNA

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The putative ABCA8 coding sequence is app. 6.5 kb in size. We successfully cloned and sequenced a 1kb segment of the human ABCA8 cDNA that encodes the putative second nucleotide binding site of the mature polypeptide (the sequence is shown in the sequence listing). The nucleotide sequence exhibits a 73% homology with the known human ABCA1 (ABC1) cDNA sequence.

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We identified an alternative transcript in the cloned 1 kb coding region which consists of a 72 bp segment (see sequence listing). Genomic analysis of this region revealed that the alternative sequence is identical with a complete intron suggesting that the alternative mRNA is generated by intron retention. The retained intron introduces a preterminal stop codon and thus may code for a truncated ABCA8 variant.

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ABCA8 also shows a cholesterol sensitive regulation of the mRNA expression (Table 2).

Tissue expression of ABCA8 is shown in table 1.

Example 9

Characterisation of the regulation of ABC transporter during differentiation of keratinocytic cells (HaCaT)

Differentiation of epidermal keratinocytes is accompanied by the synthesis of specific lipids composed mainly of sphingolipids (SL), free fatty acids (FFA), cholesterol (CH), and cholesterol sulfate, all involved in the establishment of the epidermal permeability barrier. The skin and, in particular, the proliferating layer of the epidermis is one of the most active sites of lipid synthesis in the entire organism. Cholesterol synthesis in normal human epidermis is LDL-independent, and circulating cholesterol levels do not affect the cutaneous de novo cholesterol synthesis. Fully differentiated normal human keratinocytes lack LDL receptors or its expression is very low, whereas in the normal human epidermis only basal cells express LDL receptors.

During keratinocyte differentiation a shift from polar glycerophospholipids to neutral lipids (FFA, TG) and also a replacement of short chain FFA by long chain highly saturated FFA is observed. The most important lipids for the barrier function of the skin are sphingolipids that account for one third of the lipids in the cornified layer, and consist of a large ceramide fraction as a result of glucosylceramide degradation by intercellular glycosidases and de novo synthesis of ceramide.

Glucosylceramide is synthesized intracellulary and stored in lamellar bodies and glucosylceramide synthase expression was found up-regulated during the differentiation of human keratinocytes.

Cholesterol sulfate is formed by the action of cholesterol sulfotransferase during keratinocyte differentiation. Cholesterol sulfate and the degrading enzyme steroid sulfatase are present in all viable epidermal layers, with the highest levels in the stratum granulosum. The gradient of cholesterol sulfate content across the stratum corneum (from inner to outer layers), and progressive desulfation of cholesterol sulfate regulate cell cohesiveness and normal stratum corneum keratinization and desquamation, respectively. Cholesterol sulfate induces transglutaminase 1 and the coordinate regulation of both factors is essential for normal keratinization.

The final step in lipid barrier formation involves lamellar body secretion and the subsequent post-secretory processing of polar lipids into their nonpolar lipid products through the action of hydrolytic enzymes that are simultaneously released (β-glucoccrebrosidase, phospholipases, steroid sulfatase, acid sphingomyelinase). Disruption of the permeability barrier results in an increased cholesterol, fatty acid, and ceramide synthesis in the underlying epidermis. It has been shown that mRNA levels for the key enzymes required for cholesterol, fatty acid, and ceramide synthesis increased rapidly after artificial barrier disruption.

Currently the lipid transport systems in keratinocytes are poorly characterized. Several fatty acid transport related proteins have been identified in keratinocytes: plasma membrane fatty acid transport proteins (FATP) and intracellular fatty acid binding proteins (FABPs), most of them exhibiting high affinity for essential fatty acids. The expression of epidermal FABPs is up-regulated in hyperproliferative and inflammatory skin diseases, during keratinocyte differentiation and barrier disruption

Based on our data on macrophages, we propose several ABC transporters as putative candidates for cellular lipid export in keratinocytes. We have examined the expression of all known ABC transporters during HaCaT cells differentiation. The human HaCaT cell line has a full epidermal differentiation capacity. Keratinocytes grown in

vitro as a monolayer at low calcium concentration (< 0.1 mM) can be differentiated by increasing calcium concentration in the culture medium (1-2 mM). We cultured HaCaT cells as a monolayer in calcium-free RMPI (Gibco) medium mixed with standard Ham's F12 medium at a ratio 3:1 supplemented with 10% chelex-treated FCS, Penicillin and Streptomycin. The final concentration of calcium in above medium was 0.06 mM. When the cells reached confluence (usually on 5th day of the culture), calcium concentration was enhanced up to the level of 1.2 mM. The cells were seeded at a density of 2×10^5 / cm⁻² in 60 mm culture dishes. The culture medium was replaced every two day and the cells were harvested after 24 h, 48h h, 4 d, 6 da, 8 d and 10 d in culture, respectively. Total RNA from HaCaT cells was isolated using the isothiocyanate/cesium chloride-ultracentrifugation method.

The expression of all known human ABC transporters was examined during HaCaT cell differentiation (24 h, 48 h, 4 d, 6 d, 8 d, 10d, respectively) using a semi-quantitative RT-PCR approach (Table 6). The primer sets were generated from the published sequences of the ABC-transporters. Primers specific for GAPDH were used as a control. As a marker of keratinocyte differentiation CGT (ceramide glucosyl transferase) gene expression was assessed. Three of the transporters examined, ABCB1 (MDR1), ABCB4 (MDR3), ABCD3 (PMP70), were not expressed. ABCC6 (MRP6), ABCA1 (ABC1), ABCD2 (ALDR and ABCB9 (est122234) were expressed at low levels (Table 6)

Most of the other transporters exhibited a biphasic expression pattern or were downregulated during keratinocyte differentiation. There was, however, a high expression of ABCG1 (ABC8), ABCA8 (new) and ABCC3 (MRP3) indicative for their involvement in terminal keratinocyte lipid secretion for cholesterol, FFAs and ceramide-backbone lipids.. The two peroxisomal ABC transporters, ABCD2 (ALDR) and ABCD1 (ALDP) that mediate the transport of very long chain fatty acids into peroxisomes were initially expressed at relatively low levels and subsequently downregulated during differentiation. This is in agreement with the replacement of

short chain fatty acids by very long chain fatty acids during keratinocyte differentiation.

Example 10:

Sequencing of ABCA1 cDNA and genomic structure in five families of patients with Tangier disease revealed different mutations in the ABCA1 gene locus. These patients have different mutations at different positions in the ABCA1 gene, that result in changes in the protein structure of ABCA1. Family members that are heterozygous for these mutations show lowered levels of serum HDL, whereas the homocygote patients have extremely reduced HDL serum levels.

Claims

Claims:

10		1.	A polynucleotide comprising a member selected from the group consisting of:
	5		(a) a polynucleotide encoding the polypeptide as set forth in SEQ ID NO:2;
15			(b) a polynucleotide capable of hybridizing to and which is at least 70% identical to the polynucleotide of (a); and
20	10		(c) a polynucleotide fragment of the polynucleotide of (a) or (b).
	- X	2.	The polynucleotide of claim 1 wherein the polynucleotide is DNA.
25		3.	A vector containing one or more of the polynucleotides of claim 1 and 2.
	15	4.	A host cell containing the vector of claim 3.
30		5.	A process for producing a polypeptide comprising: expressing from the host cell of claim 4 the polypeptide encoded by said DNA.
35	20	6.	A polypeptide selected from the group consisting of
40	25	·	 (a) a polypeptide having the deduced amino acid sequence of SEQ ID NO:2 and fragments, analogs and derivatives thereof, and (b) a polypeptide comprising amino acid 1 to amino acid 2201 of SEQ ID NO:2.
45		7.	An antibody capable to bind to the polypeptide of claim 6.
*	30	8.	A diagnostic kit for the detection of the polypeptide of claim 6.
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			9.	Use of a polypeptides encoded by a polynucleotide comprising a member selected from the group consisting of:
10				(a) a polynucleotide as set forth in SEQ ID NO:1, 3, 4 and 6 to 31;
	٠	5		(b) a polynucleotide capable of hybridizing to and which is at least 70% identical to the polynucleotide of (a); and
15				(c) a polynucleotide fragment of the polynucleotide of (a) or (b)
20		10		in an assay for for detecting modulators of said polypeptides.
			10.	Modulator of a polypeptides encoded by a polynucleotide comprising a member selected from the group consisting of:
25		1.6		(a) a polynucleotide as set forth in SEQ ID NO:1, 3, 4 and 6 to 31;
		15		(b) a polynucleotide capable of hybridizing to and which is at least 70% identical to the polynucleotide of (a); and
30				(d) a polynucleotide fragment of the polynucleotide of (a) or (b)
0.5		20	11.	A pharmaceutical comprising the modulator of claim 10
35			12.	An assay for detecting polypeptides encoded by a polynucleotide comprising
			•	a member selected from the group consisting of:
40				(a) a polynucleotide as set forth in SEQ ID NO:1, 3, 4 and 6 to 32 and 54;
		25		(b) a polynucleotide capable of hybridizing to and which is at least 70% identical to the polynucleotide of (a); and
45				(c) a polynucleotide fragment of the polynucleotide of (a) or (b)

Figure 1

Figure 2

1 CAAACATGTCAGCTGTTACTGGAAGTGGCCTGGCCTCTATTTATCTTCCTGATCCTGATC 61 TCTGTTCGGCTGAGCTACCCACCCTATGAACAACATGAATGCCATTTTCCAAATAAAGCC 120 121 ATGCCCTCTGCAGGAACACTTCCTTGGGTTCAGGGGATTATCTGTAATGCCAACAACCCC 180 1 M P S A G T L P W V Q G I I C N A N N P 181 TGTTTCCGTTACCCGACTCCTGGGGAGGCTCCCGGAGTTGTTGGAAACTTTAACAAATCC 240 21 C F R Y P T P G E A P G V V G N F N K S 241 ATTGTGGCTCGCCTGTTCTCAGATGCTCGGAGGCTTCTTTTATACAGCCAGAAAGACACC 300 41 I V A R L F S D A R R L L L Y S Q K D T 60 301 AGCATGAAGGACATGCGCAAAGTTCTGAGAACATTACAGCAGATCAAGAAATCCAGCTCA 360 61 S M K D M R K V L R T L Q Q I K K S S S 361 AACTTGAAGCTTCAAGATTTCCTGGTGGACAATGAAACCTTCTCTGGGTTCCTGTATCAC 420 81 N L K L Q D F L V D N E T F S G F L Y H 421 AACCTCTCTCCCAAAGTCTACTGTGGACAAGATGCTGAGGGCTGATGTCATTCTCCAC 480 101 N L S L P K S T V D K M L R A D V I L H 120 481 AAGGTATTTTTGCAAGGCTACCAGTTACATTTGACAAGTCTGTGCAATGGATCAAAATCA 540 121 K V F L Q G Y Q L H L T S L C N G S K S 541 GAAGAGATGATTCAACTTGGTGACCAAGAAGTTTCTGAGCTTTGTGGCCTACCAAGGGAG 600 141 E E M I Q L G D Q E V S E L C G L P R E 601 AAACTGGCTGCAGCAGAGCGAGTACTTCGTTCCAACATGGACATCCTGAAGCCAATCCTG 660 161 K L A A A E R V L R S N M D I L K P I L 661 AGAACACTAAACTCTACATCTCCCTTCCCGAGCAAGGAGCTGGCCGAAGCCACAAAAACA 720 181 R T L N S T S P F P S K E L A E A T K T 200 721 TTGCTGCATAGTCTTGGGACTCTGGCCCAGGAGCTGTTCAGCATGAGAAGCTGGAGTGAC 780 201 L L H S L G T L A Q E L F S M R S W S D 220 781 ATGCGACAGGAGGTGATGTTTCTGACCAATGTGAACAGCTCCAGCTCCTCCACCCAAATC 840 221 M R Q E V M F L T N V N S S S S T Q I 241 Y Q A V S R I V C G H P E G G G L K I K 260 901 TCTCTCAACTGGTATGAGGACAACAACTACAAAGCCCCTCTTTGGAGGCAATGGCACTGAG 960 261 S L N W Y E D N N Y K A L F G G N G T E

961 GAAGATGCTGAAACCTTCTATGACAACTCCTACAACTCCTTACTGCAATGATTTGATGAAG 1020 281 E D A E T F Y D N S T T P Y C N D L M K 300 1021 AATTTGGAGTCTAGTCCTCTTTCCCGCATTATCTGGAAAGCTCTGAAGCCGCTGCTCGTT 1080 301 N L E S S P L S R I I W K A L K P L L V 1081 GGGAAGATCCTGTATACACCTGACACTCCAGCCACAAGGCAGGTCATGGCTGAGGTGAAC 1140 321 G K I L Y T P D T P A T R Q V M A E V N 1141 AAGACCTTCCAGGAACTGGCTGTTTCCATGATCTGGAAGGCATGTGGGAAGTCAGC 1200 341 K T F Q E L A V F H D L E G M W E E L S 360 1201 CCCAAGATCTGGACCTTCATGGAGAACAGCCAAGAAATGGACCTTGTCCGGATGCTGTTG 1260 361 P K I W T F M E N S Q E M D L V R M L L 1261 GACAGCAGGGACAATGACCACTTTTGGGAACAGCAGTTGGATGGCTTAGATTGGACAGCC 1320 381 D S R D N D H F W E Q Q L D G L D W T A 1321 CAAGACATCGTGGCGTTTTTGGCCAAGCACCCAGAGGATGTCCAGTCCAGTAATGGTTCT 1380 401 Q D I V A F L A K H P E D V Q S S N G S 1381 GTGTACACCTGGAGAGAGCTTTCAACGAGACTAACCAGGCAATCCGGACCATATCTCGC 1440 421 V Y T W R E A F N E T N Q A I R T I S R 1441 TTCATGGAGTGTGTCAACCTGAACAAGCTAGAACCCATAGCAACAGAAGTCTGGCTCATC 1500 441 F M E C V N L N K L E P I A T E V W L I 1501 AACAAGTCCATGGAGCTGCTGGATGAGAGGAAGTTCTGGGCTGGTATTGTGTTCACTGGA 1560 461 N K S M E L L D E R K F W A G I V F T G 480 1561 ATTACTCCAGGCAGCATTGAGCTGCCCCATCATGTCAAGTACAAGATCCGAATGGACATT 1620 481 I T P G S I E L P H H V K Y K I R M D I 1621 GACAATGTGGAGAGGACAAATAAAATCAAGGATGGGTACTGGGACCCTGGTCCTCGAGCT 1680 501 D N V E R T N K I K D G Y W D P G P R A 1681 GACCCCTTTGAGGACATGCGGTACGTCTGGGGGGGGGCTTCGCCTACTTGCAGGATGTGGTG 1740 521 D P F E D M R Y V W G G F A Y L Q D V V 1741 GAGCAGCCAATCATCAGGGTGCTGACGGGCACCGAGAAGAAAACTGGTGTCTATATGCAA 1800 541 E Q A I I R V L T G T E K K T G V Y M Q 1801 CAGATGCCCTATCCCTGTTACGTTGATGACATCTTTCTGCGGGTGATGAGCCGGTCAATG 1860 561 Q M P Y P C Y V D D I F L R V M S R S M 1861 CCCCTCTTCATGACGCTGGCCTGGATTTACTCAGTGGCTGTGATCATCAAGGGCATCGTG 1920 581 P L F M T L A W I Y S V A V I I K G I V 1921 TATGAGAAGGAGGCACGGCTGAAAGAGACCATGCGGATCATGGGCCCTGGACAACAGCATC 1980 601 Y E K E A R L K E T M R I M G L D N S I 1981 CTCTGGTTTAGCTGGTTCATTAGTAGCCTCATTCCTCTTCTTGTGAGCGCTGGCCTGCTA 2040 621 L W F S W F I S S L I P L L V S A G L L 2041 GTGGTCATCCTGAAGTTAGGAAACCTGCTGCCCTACAGTGATCCCAGCGTGGTGTTTGTC 2100 641 V V I L K L G N L L P Y S D P S V V F V

2101 TTCCTGTCCGTGTTGCTGGTGACAATCCTGCAGTGCTTCCTGATTAGCACACTCTTC 2160

661 F L S V F A V V T I L Q C F L I S T L F 2161 TCCAGAGCCAACCTGGCAGCAGCCTGTGGGGGCATCATCTACTTCACGCTGTACCTGCCC 2220 681 S R A N L A A A C G G I I Y F T L Y L P 2221 TACGTCCTGTGTGGCATGGCAGGACTACGTGGGCTTCACACTCAAGATCTTCGCTAGC 2280 701 Y V L C V A W Q D Y V G F T L K T F A S 2281 CTGCTGTCTCCTGTGGCTTTTGGGTTTGGCTGTGAGTACTTTGCCCTTTTTGAGGAGCAG 2340 721 L L S P V A F G F G C E Y F A L F E E Q 2341 GGCATTGGAGTGCAGTGGGACAACCTGTTTGAGAGTCCTGTGGAGGAAGATGGCTTCAAT 2400 741 G I G V Q W D N L F E S P V E E D G F N 2401 CTCACCACTTCGGTCTCCATGATGCTGTTTGACACCTTCCTCTATGGGGTGATGACCTGG 2460 761 L T T S V S M M L F D T F L Y G V M T W 2461 TACATTGAGGCTGTCTTTCCAGGCCAGTACGGAATTCCCAGGCCCTGGTATTTTCCTTGC 2520 781 Y I E A V F P G Q Y G I P R P W Y F P C 800 2521 ACCAAGTCCTACTGGTTTGGCGAGGAAAGTGATGAGAAGAGCCCACCCTGGTTCCAACCAG 2580 801 T K S Y W F G E E S D E K S H P G S N Q 821 K R I S E I C M E E E P T H L K L G V S 2641 ATTCAGAACCTGGTAAAAGTCTACCGAGATGGGATGAAGGTGGCTGTCGATGGCCTGGCA 2700 841 I Q N L V K V Y R D G M K V A V D G L A 2701 CTGAATTTTTATGAGGGCCAGATCACCTCCTTCCTGGGCCACAATGGAGCGGGGAAGACG 2760 861 L N F Y E G Q I T S F L G H N G A G K T 2761 ACCACCATGTCAATCCTGACCGGGTTGTTCCCCCCGACCTCGGGCACCGCCTACATCCTG 2820 881 T T M S I L T G L F P P T S G T A Y I L 2821 GGAAAAGACATTCGCTCTGAGATGAGCACCATCCGGCAGAACCTGGGGGTCTGTCCCCAG 2880 901 G K D I R S E M S T I R Q N L G V C P Q 2881 CATAACGTGCTGTTTGACATGCTGACTGTCGAAGAACACATCTGGTTCTATGCCCGCTTG 2940 921 H N V L F D M L T V E E H I W F Y A R L 2941 AAAGGGCTCTCTGAGAAGCACGTGAAGGCGGAGATGGACGAGATGGCCCTGGATGTTGGT 3000 S E K H V K A E M E Q M A L D V G 3001 TTGCCATCAAGCAAGCTGAAAAGCAAAACAAGCCAGCTGTCAGGTGGAATGCAGAGAAAG 3060 961 L P S S K L K S K T S Q L S G G M Q R K 980 3061 CTATCTGTGCCCTTGGCCTTTGTCGGGGGATCTAAGGTTGTCATTCTGGATGAACCCACA 3120 981 L S V A L A F V G G S K V V I L D E P T 3121 GCTGGTGTGGACCCTTACTCCCGCAGGGGAATATGGGAGCTGCTGCTGAAATACCGACAA 3180 1001 A G V D P Y S R R G I W E L L L K Y R Q 3181 GGCCGCACCATTATTCTCTCTACACACCACATGGATGAAGCGGACGTCCTGGGGGACAGG 3240 1021 G R T I I L S T H H M D E A D V L G D R 1040 3241 ATTGCCATCATCTCCCATGGGAAGCTGTGCTGTGTGGGCTCCTCCTGTTTCTGAAGAAC 3300 1041 I A I I S H G K L C C V G S S L F L K N

1061 Q L G T G Y Y L T L V K K D V E S S L S 1080 3361 TCCTGCAGAAACAGTAGCACTGTGTCATACCTGAAAAAGGAGGACAGTGTTTCTCAG 3420 1081 S C R N S S S T V S Y L K K E D S V S Q 1100 3421 AGCAGTTCTGATGCTGGCCTGGGCAGCGACCATGAGAGTGACACGCTGACCATCGATGTC 3480 1101 S S S D A G L G S D H E S D T L T I D V 3481 TCTGCTATCTCCAACCTCATCAGGAAGCATGTGTCTGAAGCCCGGCTGGTGGAAGACATA 3540 1121 S A I S N L I R K H V S E A R L V E D I 1140 1141 G H E L T Y V L P Y E A A K E G A F V E 3601 CTCTTTCATGAGATTGATGACCGGCTCTCAGACCTGGGCATTTCTAGTTATGGCATCTCA 3660 1161 L F H E I D D R L S D L G I S S Y G I S 1180 3661 GAGACGACCCTGGAAGAATATTCCTCAAGGTGGCCGAAGAGAGTGGGGTGGATGCTGAG 3720 1181 E T T L E E I F L K V A E E S G V D A E 3721 ACCTCAGATGGTACCTTGCCAGCAAGACGAAACAGGCGGGCCTTCGGGGACAAGCAGAGC 3780 1201 T S D G T L P A R R N R R A F G D K Q S 1220 3781 TGTCTTCGCCCGTTCACTGAAGATGATGCTGCTGATCCAAATGATTCTGACATAGACCCA 3840 1221 C L R P F T E D D A A D P N D S D I D P 3841 GAATCCAGAGAGACAGACTTGCTCAGTGGGATGGCAAAGGGTCCTACCAGGTGAAA 3900 1241 E S R E T D L L S G M D G K G S Y Q V K 1260 3901 GGCTGGAAACTTACACAGCAACAGTTTGTGGCCCCTTTTGTGGAAGAGACTGCTAATTGCC 3960 1261 G W K L T Q Q Q F V A L L W K R L L I A 1280 3961 AGACGGAGTCGGAAAGGATTTTTTGCTCAGATTGTCTTGCCAGCTGTGTTTTGTCTGCATT 4020 1281 R R S R K G F F A Q I V L P A V F V C I 1300 4021 GCCCTTGTGTTCAGCCTGATCGTGCCACCCTTTGGCAAGTACCCCAGCCTGGAACTTCAG 4080 1301 A L V F S L I V P P F G K Y P S L E L Q 1320 4081 CCCTGGATGTACAACGAACAGTACACATTTGTCAGCAATGATGCTCCTGAGGACACGGGA 4140 1321 P W M Y N E Q Y T F V S N D A P E D T G 4141 ACCCTGGAACTCTTAAACGCCCTCACCAAAGACCCTGGCTTCGGGACCCGCTGTATGGAA 4200 1341 T L E L L N A L T K D P G F G T R C M E 1360 4201 GGAAACCCAATCCCAGACACGCCCTGCCAGGCAGGGAGGAAGAGTGGACCACTGCCCCA 4260 1361 G N P I P D T P C Q A G E E W T T A P 1380 4261 GTTCCCCAGACCATCATGGACCTCTTCCAGAATGGGAACTGGACAATGCAGAACCCTTCA 4320 1381 V P Q T I M D L F Q N G N W T M Q N P S 4321 CCTGCATGCCAGTGTAGCAGCGACAAAATCAAGAAGATGCTGCCTGTGTGTCCCCCAGGG 4380 1401 P A C Q C S S D K I K K M L P V C P P G 4381 GCAGGGGGGCTGCCTCCACAAAGAAAACAAAACACTGCAGATATCCTTCAGGACCTG 4440 1421 A G G L P P P Q R K Q N T A D I L Q D L 1440 4441 ACAGGAAGAAACATTTCGGATTATCTGGTGAAGACGTATGTGCAGATCATAGCCAAAAGC 4500 1441 T G R N I S D Y L V K T Y V Q I I A K S 4501 TTAAAGAACAAGATCTGGGTGAATGAGTTTAGGTATGGCGGCTTTTCCCTGGGTGTCAGT 4560

1461 L K N K I W V N E F R Y G G F S L G V S 4561 AATACTCAAGCACTTCCTCCGAGTCAAGAAGTTAATGATGCCACCAAACAATGAAGAAA 4620 1481 N T Q A L P P S Q E V N D A T K Q M K K 1500 4621 CACCTAAAGCTGGCCAAGGACAGTTCTGCAGATCGATTTCTCAACAGCTTGGGAAGATTT 4680 1501 H L K L A K D S S A D R F L N S L G R F 1520 4681 ATGACAGGACTGGACACCAGAAATAATGTCAAGGTGTGGTTCAATAACAAGGGCTGGCAT 4770 1521 M T G L D T R N N V K V W F N N K G W H 1540 4741 GCAATCAGCTCTTTCCTGAATGTCATCAACAATGCCATTCTCCGGGCCCAACCTGCAAAAG 4800 1541 A I S S F L N V I N N A I L R A N L Q K 1560 4801 GGAGAGAACCCTAGCCATTATGGAATTACTGCTTTCAATCATCCCCTGAATCTCACCAAG 4860 1561 G E N P S H Y G I T A F N H P L N L T K 1580 4861 CAGCAGCTCTCAGAGGTGGCTCCGATGACCACATCAGTGGATGTCCTTGTGTCCATCTGT 4920 1581 Q Q L S E V A P M T T S V D V L V S I C 1600 4921 GTCATCTTTGCAATGTCCTTCGTCCCAGCCAGCTTTGTCGTATTCCTGATCCAGCAGCGG 4980 1601 V I F A M S F V P A S F V V F L I Q E R 1620 4981 GTCAGCAAAGCAAAACACCTGCAGTTCATCAGTGGAGTGAAGCCTGTCATCTACTGGCTC 5040 1621 V S K A K H L Q F I S G V K P V I Y W L 1640 5041 TCTAATTTTGTCTGGGATATGTGCAATTACGTTGTCCCTGCCACACTGGTCATTATCATC 5100 1641 S N F V W D M C N Y V V P A T L V I I I 5101 TTCATCTGCTTCCAGCAGAAGTCCTATGTGTCCTCCACCAATCTGCCTGTGCTAGCCCTT 5160 1661 F I C F Q Q K S Y V S S T N L P V L A L 1680 5161 CTACTTTTGCTGTATGGGTGGTCAATCACACCTCTCATGTACCCAGCCTCCTTTGTGTTC 5220 1681 L L L Y G W S I T P L M Y P A S F V F 1700 5221 AAGATCCCCAGCACAGCCTATGTGGTGCTCACCAGCGTGAACCTCTTCATTGGCATTAAT 5280 1701 K I P S T A Y V V L T S V N L F I G I N 1720 5281 GGCAGCGTGGCCACCTTTGTGCTGGAGCTGTTCACCGACAATAAGCTGAATAATATCAAT 5340 1721 G S V A T F V L E L F T D N K L N N I N 1740 5341 GATATCCTGAAGTCCGTGTTCTTGATCTTCCCACATTTTTGCCTGGGACGAGGGCTCATC 5400 1741 DILKSVFLIFPHFCLGRGLI 1760 5401 GACATGGTGAAAAACCAGGCAATGGCTGATGCCCTGGAAAGGTTTGGGGGAGAATCGCTTT 5460 1761 D M V K N Q A M A D A L E R F G E N R F 1780 5461 GTGTCACCATTATCTTGGGACTTGGTGGGACGAAACCTCTTCGCCATGGCCGTGGAAGGG 5520 1781 V S P L S W D L V G R N L F A M A V E G 1800 5521 GTGGTGTTCTTCCTCATTACTGTTCTGATCCAGTACAGATTCTTCATCAGGCCCAGACCT 5580 1801 V V F F L I T V L I Q Y R F F I R P R P 1820 5581 GTAAATGCAAAGCTATCTCCTCTGAATGATGAAGATGAAGATGTGAGGCGGGAAAGACAG 5640 1821 V N A K L S P L N D E D E D V R R E R Q 1840 5641 AGAATTCTTGATGGTGGAGGCCAGAATGACATCTTAGAAATCAAGGAGTTGACGAAGATA 5700 1841 R I L D G G G Q N D I L E I K E L T K I 1860 5701 TATAGAAGGAAGCCGGAAGCCTGCTGTTGACAGGATTTCCGTGGGGCATTCCTCCTGGTGAG 5760

1861		R	R	K	R	K	₽	A	V	D	R	I	C	V		I	P	P	G	E	1880
5761	TG	CTT	TGG	GCI	CCI	GGG	AGT	TAA	TGG	GGC	TGG	AAA	ATC	ATC	AAC	TTT	'CAA	GAT	GTT	AACA	5820
1881		F	G	L	L	G	v	N	G	A	G	K	s	s	T	F	K	M	L	T	1900
5821	GG	AGA	TAC	CAC	TGT	TAC	CAG	AGG	AGA	TGC	TTT	CCI	TAA	CAG	AAA	TAC	TAT	CTT	ATC	AAAC	5880
1901	G	D	T	T	V	T	R	G	D	A	F	L	N	R	N	s	I	L	s	N	1920
5881	ΑT	CCA	TGA	AGT	ACA	TCA	GAA	CAT	GGG	CTA	CTG	CCC	TCA	GTT	TGA	TGC	CAT	CAC	AGA	GCTG	5940
1921	I	н	E	v	Н	Q	N	М	G	Y	С	P	Q	F	D	A	I,	т	E	L ·	1940
5941	TT	GAC	TGG	GAG	AGA	ACA	CGI	GGA	GTI	CTT	TGC	CCI	TTT	GAG	AGG	AGI	ccc	AGA	GAA.	AGAA	6000
1941	<u>r</u>	T	G	R	E	н	v	E	F	F	A	L	L	R	G	v	P	E	к	E	1960
6001	GI	TGG	CAA	GGI	TGG	TGA	GTG.	GGC	GAT	TCG	GAA	ACI	GGG	CCT	CGI	GAA	GTA	TGG	AGA	AAAA	6060
1961	<u>v</u>	G	ĸ	v	G	E	W	Α	I	R	K	L	G	L	v	к	Y	G	E	к	1980
6061	TA	TGC	TGG	TAA	CTA	TAG	TGG	AGG	CAA	CAA	ACG	CAA	GCT	CTC	TAC	AGC	CAT	GGC	TTT	GATC	6120
1981	Y	A	G	N	Y	s	G	G	N	ĸ	R	ĸ	L	s	T	A	M	A	L	ī.	2000
6121	GG	CGG	GCC	TCC	TGT	GGT	GTT	TCI	GGA	TGA	ACC	CAC	CAC	AGG	CAT	GGA	TCC	CAA	AGC	CCGG	6180
2001	G	_G	P	P	v	v	F	L	D	E	P	T	T	G	м	ם	P	к	A	R	2020
6181	CG	GTT	CTT	GTG	GAA	TTG	TGC	CCI	AAG	TGT	TGT	CAA	GGA	GGG	— GAG	ATC	AGT	AGT	GCT	TACA	6240
2021		F	L	W	N	С	A	T.	s	v	v	к	E	G	R	s	v	v	L	T	2040
6241	TC	TCA	TAG	TAT	CGA	AGA	ATG	TGA	AGC	TCT	TTG	CAC	TAG	GAT	GGC	:AAI	CAT	GGT	CAA	TGGA	6300
2041	s	н	s	M	E	E	С	E	A	L	С	T	R	м	A	I	M	v	N	G	2060
6301	AG	GTT	CAG	GTG	CCT	TGG	CAG	TGT	CCA	GCA	TCT	AAA	AAA	TAG	GTT	TGG	AGA	TGG	TTA	TACA	6360
2061	R	F	R	С	L	G	s	v	Q	н	L	ĸ	N	R	F	G	ם	G	Y	T	2080
6361	AT	AGT	TGT	ACG	AAT	AGC	AGG	GTC	CAA	ccc	GGA	CCI	GAA	GCC	TGT	CCA	GGA	TTT	CTT	TGGA	6420
2081		v	v	R	I	A	G	s	N	P	D	L	к	p	v	Q	ם	F	F	G	2100
6421	CI	TGC	ATT	TCC	TGG	AAG	TGT	TCC	AAA	AGA	GAA	ACA	.CCG	GAA	CAT		ACA	ATA	CCA	GCTT	6480
2101		A	F	P	G	s	v.	P	к	E	к	н	R	N	м	L	o	Y	Q	L	2120
6481	CC	ATC	TTC	ATI	ATC	TTC	TCT	GGC	CAG	GAT	ATT	CAG	CAT	CCT	CTC	CCA	.GAG		_	GCGA	
2121		s	s	L	s	s	L	A	R	ľ	F	s	Í	L	s	Q	s	к	ĸ	R	2140
6541	CI	CCA	CAT	AGA	AGA	CTA	CTC	TGT	TTC	TCA	GAC	AAC	ACT	TGA	CĊA	AGI	ATT	TCT	GAA	CTTT	6600
2141		н	I	E	D	Y	s	v	s	o	T	Ŧ	L	D		v	F	v	N	F	2160
6601	GC	CAA	CCA	CCA	AAG	TGA	TGA	TGA	CCA	CTT	AAA	AGA	CCT	CTC	_		CAA	AAA	CCA	GACA	
2161		ĸ	α	Q	s	D	D	מ	н	L	ĸ	D	L	s	L	н	к	N	Q	T	2180
6661	GI	'AGT	GGA	CGI	TGC	AGT	TCT	CAC	ATC	TTT	TCT	ACA	GGA	TGA	GAA				_		6720
2181		v	D	v	A	v	L	T	s	F	L	Q	D	E	ĸ	v	к	E	s	Y	2200
6721	GI	'ATG	AAG	AAT	CCI	GTT	CAT					_							_		6780
2201		*																			00
6781	GC	ACC	ATC	TGA	AGT	GTT	GTC	GAC	AAA	ACA	GCC	AGA	AGT	TGA	TGT	'GCC	AAC	מאג	AAT	ACTG	6840
6841																					6880
_	_	_																			

Figure 3

5' 1 GTACCCCCT TGCCTGGTTG ATCCTCAGGG TTCTACTTAG AATGCCTCGA

51	AAAGTCTTGG	CTGGACACCC	ATGCCCAGTC	TTTCTGCAGG	GTCCCATTGG
101	GGTTAACCTT	CTCATTTCAT	CCCATGTGAA	CCAGGCCAGG	CCCATCAGGG
151	TTTGGCAACC	CCCTGATGCA	GTGGTTGCTG	CCAGGTGACA	GGAGCAAGCC
201	TGCAGCTGCT	GGGGGCCAT	GCAGAGACAG	CCTGCCAGAG	GGGAGACCAC
251	CTGGGGAGGC	CAGAGCCGTG	GAGACAGCAA	GAGACCAGGG	GCTGAGGACA
301	GAGTAGTACA	${\tt GGTCTTTGGT}$	CCCAGTAGTC	CTGAAACCAC	TGCACTCCGA
351	ACCTTTCTGT	ACTTAGCTTA	AGCCAGTTGG	AGTTTCTGTC	CTTTACAACC
401	AAGAGCCTTG	ATAGGAATGG	GGTCCTGTGC	TACGCTACTG	TTGGCTTCTT
451	TCCCGATCGG	GCGCTGGAGG	GGAACACAGC	AGTGACTACA	GTGGGATGCT
501	TACTCGGTGC	TGGGCATGCT	AGAAAGTGCT	TGCCATGCCT	TATTTCCCAC
551	GTGGTGGGGA	TTTTGACCCC	ACCTGTACAG	ACAGATAAGT	GAGGACCCTT
601	TTCACCTTAT	CCTGCAACAG	AAAATCCAGC	AGCCAAAGCC	AACAAGGGCC
651	CAGCATAGCA	TCTTCCCTCT	CTGACTTCAT	CCTCACGCTC	CACACACCAT
701	CCCCCTGGCC	ATTCCCAGCA	GCCCAGTAAG	CACTGCCTCA	CACTTCCAGT
751	TCCGGACCAG	CCAGGATGCC	CAGGCTGGAT	GGGGGCCATC	CACCGGCTGA
801	AGCCAATTGC	$\mathtt{CTATTCTCGA}$	GCTGAAGGTG	AATCAATCCC	GCATAAATCT
851	TCGGGCAGAG	AACTNGGGTG	GGGGGTAGAA	GAGGGGGAAT	GTCTAGAAGG
901	AAATTCTGGG	${\tt GCACATTCCT}$	GGAAGTGAGG	AGGATGGATA	TTGGACAGAA
951	ATTATGTCAT	TGCAGGCACC	CTCACTTGCC	CTGGCCACAT	GGACAGTTCC
1001	TCCCCGGCTG	TGTTCCGNGC	CTCCTCTCGT	GCTCCAGGGC	CTGTCTGTTC
1051	CTGGAGCGAG	ATGGGTCCCA	GGGCTGGGCA	CCAGTCCCCA	TCTCCAGCCA
1101	TCAGGCACTT	TCCTCTCTGT	GTTTTGGCGT	AAACACNTCC	CTAGGTTTGT
1151	GGATCTGAAT	CCTCTTCCCA	ACACACTCAA	GCTTTGCTGG	GCCTCCCTGC
1201	AGTGTATGTT	TAAGGCACCA	CACAGCCTCC	AAGGCCTGGC	ACCCGGGCAG
1251	TGGCCACCTG	GTAAACACAG	CAGTCAGATT	TCCTCATTTC	AGCCAAGTGT
1301	AAAATCAAGG	TAATGGATCT	ACNCTTTTTT	TTTTNTNTTT	TTTCCAGGG
1351,	GNTNNTTTTT	TTTTGAGACG	GAGTCTCACT	CTGTCANCCC	CGGTCTGGAG
1401	TGCAGTGGCT	CAATCTCGGC	TCANCTGGCA	AGCTCCGCCT	CCCAGGTTCA
1451	TGCCATTCTC	CTGCCTCAGC	CTACATAGTA	GCTGGGACTA	CAGGTGCCCG
1501	CCACCACACC	TAGCTAATTT	TTTGTATTTT	TAGTAGAGAC	GGGGTTTCAT
1551	CATGTTAGCC	AGGATGGTCT	CGATCTCCTG	ACCTCCCAAA	GTGGTGGGAG
1601	TTACAGGTGT	GAGCCACTGC	GCNCCGGCTG	GATGACTCTT	GAGACAACAC
1651	CATTCAGACA	AAGGCAAGGC	CTCCCACTTA	AACTCATAAC	CGTGTCTCCT
1701	TTCTCTCCTT	CGATTTGAGC	GGCTGAATTT	GGTTACAGTC	ATCTGACCTG
1751	TGGGTGTGAA	NGTCCACCTG	CCTGGCATAA	AAAGCTGTGC	CTCCTTTCTA
1801	GGTGAGGAGA	AAGAGAGAGA	CCTGGCTCAT	CTGAGGTGTG	GTTGGGAGGG
1851	GGGACCCAGG	TGTGCTGGAA	ATGAAAAGAA	ATGCATTCCT	GTTTTTTCGT
1901	CCCAACATGC	AAACAACTGA	ACAAAAGCAT	TAGGGCCTGA	GACTGGGAGT
1951	AAAGAATTCC	TTGTCACCAT	GGATACCAGG	AAATGGCCCC	ACTINTATAT
2001	AATAAGGGCT	TTAGAGATGC	TGGACCATCT	GATATTCCAG	CCTGGGGCCA
2051	CATGGGAGTG	TGCCCTGGTG	TTATTCCTTA	TACAGTTCCA	TGAACATGGC
2101	TCTGGAAACA	CCTCTGTCTG	CAGAAAATGA	GGCTTTTCTT	TTTTTGTTCG

2151	GGGGTGAACA	GAGGGCAGAG	GCCTGGGCAT	CTTCACTCAG	CACCCCTTTG
2201	TAACCCAGCA	CTTAGCACCA	TGGCTGGCGC	ACAGCAATGT	CACATGTGTG
2251	AGTGCACACG	ATGCCTCACT	GCCAGGGGTC	ACCCCACACC	GGTGCTGTTG
2301	GGGGCGTTGG	AGTGGTTATC	TCTTCTTTAG	TCCTCAAGCT	CCTACCTGGC
2351	AGAGAGCTGC	CCAACACCGT	CGGGGTGGGG	TGGGCGGGAA	GGGAAGAAGC
2401	AGCAGCAAGA	AAGAAGCCCC	CTGGCCCTCA	CTCTCCCTCC	CTGGACGCCC
2451	CCTCTTCGAC	CCCATCACAC	AGCCGCTTGA	GCCTTGGAGN	CAGTGGATTT
2501	CCGAGCCTGG	GAACCCCCGG	CGTCTGTCCC	GGTGTCCCCC	GCAGCCTCAC
2551	CCNCGTGCTG	GCCCAGCCCC	CGCGAGTTCG	GGACCCGGGG	TTTCCGGGGT
2601	GGCAGGGGGT	TCCCATGCCG	CCTGCGAGGC	CTCGGCTCGG	GCCGCTCCCG
2651				GCCCCCAGCA	
2701				CCCCTTGGTG	
2751				AACCAGAGCC	
2801				CCGCGCAGCG	
2851				AGCCTCGTCC	
2901	CCGCCGCACG	CCGCCGCCGC	CCCCCCGGG	GCATGGCTGT	CTGATGGCCG
			CON1/INTRON	_	
2951				AGCGCATCCT	
3001				AACACACAAA	
3051				CCTCTGGGGC	
3101				CGCTGCCCCT	
3151				CCTTGCACCG	
3201				CGGGGGGCAG	
3251				GGCTTCTGGG	
3301				ACTGGGGAGG	
3351				GGGGAGCCCC	
3401				GCCATCCCCA	
3451			GAGGAAGCTG	CCCCCAGAGA	GCCGGAGCTC
3501	GACTGNACTC	CC 3'			

Figure 4

5′

1 CTTGGTGCCG CATGCATCGT GGTGCTCATC TTTCTGGCCT TCCAGCAGAG
51 GGCATATGTG GCCCCTGCCA ACCTGCCTGC TCTCCTGCTG TTGCTACTAC
101 TGTATGGCTG GTCGATCACA CCGCTCATGT ACCCAGCCTC CTTCTTCTTC
151 TCCGTGCCCA GCACAGCCTA TGTGGTGCTC ACCTGCATAA ACCTCTTTAT
201 TGGCATCAAT GGAAGCATGG CCACCTTTGT GCTTGAGCTC TTCTCTGATC
251 AGAAGCTGCA GGAGGTGAGC CGGATCTTGA AACAGGTCTT CCTTATCTTC
301 CCCACTTCTG CTTGGGCCGG GGGCTTATTG ACATGGTGCG GNAACCAGGC
351 CATGGCTGAT GCCTTTGANC CCTTGGGAAA AAGGCAGTTC AAGTACCCTG

401	ncttggaagg	TGGCGGAAGA	ACCTTTTGGC	ATGGGAACAG	GGCCCCTTTT
451	CCTTCTCTTC	ACACTANTGT	TCAAGCACCG	AAGCCAACTC	NTGCCACAAG
501	CCCAGGTAAG	GTCTCTGCCA	CTCCTGGAGA	GAGACGAGGA	TGTAGCCCGT
551	GAACGGGAGC	GGGTGGTCCA	AGGAGCCACC	CAGGGGGATG	TGTTGGTGCT
601	GAGGAACTTG	ACCAAGGTAT	ACCGTGGGCA	GAGGATGCCA	GCTGTTGACC
651	GCTTGTGCCT	GGGGATTCCC	CCTGGTGAGT	GTTTTGGGCT	GCTGGGTGTG
701	AACGGAGCAG	GGAAGACGTC	CACGTTTCGC	ATGGTGACGG	GGGACACATT
751	GGCCAGCAGG	GGCGAGGCTG	TGCTGGCAGG	CCACAGCGGG	CCCGGGAACC
801	CAGTGTGCGC	ACCTCNAGGG	CAGGCNCAGC	GTGGCCCGGG	AACCCAGTGC
851	TGCGCACCTA	AGCATGGGAT	ACTGCCCTNA	ATCCGATGCC	ATCTTTGAGC
901	TGCTGACGGG	CCGCGAGCAC	CTGGAGCTGC	TTGCGCGCCT	GCGCGGTGTC
951	CCGGAGGCCC	AGGTTGCCCA	NACCGNTGGC	TCGGGCCTGG	CGCGTCTGGG
1001	ACTCTCATGG	TACGCAGACC	GGCCTGCAGG	CACCTACAGG	AACCTGCCCG
1051	GGCGGCCGCT	CGAGCCCNTA	NNTGAAGTA	31	

Figure 4b

...CTCCTGCCAC AGTTAGTGAG GTCTATGGAG AGGGTGGCAG GGGCCAAGGA
CCTACTTTAA GCCCACAGAT ATTCTGTCCC CAGGCCCAGG GTGAGGTCTC...

Figure 5

CDNA-sequences of lipid sensitive Genes:
ABCB9, ABCA6, ABCC4, ABCA1, ABCD2, ABCB1, ABCB4, ABCC2, ABCD1, ABCC1,
ABCB6, ABCB11, ABCG2, ABCC5, ABCC5, ABCG1, ABCA3

ABCB9 GENBANK: U66676

GCCAATGNCACGGTTTCATCATGGAACTCCAGGACGGCTACAGCACAGAGACAGGGGAGA AGGGCGCCAGCTGTCAGGTGGCCAGAAGCAGCGGTGGCCATGGCCGNGGCTCTGGTGC GGAACCCCCAGTCCTCATCCTGGATGAAGCCACCAGCGCTTTGGATGCCGAGAGCGAGT ATCTGATCCAGCAGGCCATCCATGGCAACCTGTCAGAAGCACACGGTACTCATCATCGCG CACCGGCTGAGCACCGTGGAGCACGCGCACCTCATTGTGGTGCTGGACAAGGGCCGCGTA GTTGCAGCGGCAGATGTGGGGTTTCAAGGCCGCAGACTTCACAGCTGGCCACAACGAGCC TGTAGCCAACGGGTCACAAGGCCTGATGGGGGGCCCCTCCTTCGCCCGGTGGCAGAGGAC CCGGTGCCTGCCTGGCAGATGTGCCCACGGAGGTTTCCAGCTGCCCTACCGAGCCCAGGC $\tt CTGCAGCACTGAAAGACGACCTGCCATGTCCCATGATCACCGCTTNTGCAATCTTGCCCC$ TGGTCCCTGCCCCATTCCCAGGGCACTCTTACCCCNNNCTGGGGGATGTCCAAGAGCATACGGGATTTTCCGTCTCTCCCTCTTGCCAGCTCTGTGAGTCTGGCCAGGGCGGGTAGGGAG CGTGGAGGGCATCTGTCTGCCAATTGCCCGCTGCCAATCTAAGCCAGTCTCACTGTGACCACACGAAACCTCAACTGGGGGGTGAGGAGCTGGCCAGGTCTGGAGGGGCCTCAGGTGCC CCCACACCCGCCCTGTGCTCTGCTGTCTGGAGGCCACGTGGACCTTCATGAGATGCATTCTCTTCTGTCTTTGGTGGANGGGATGGTGCAAAGCCCAGGATCTGGCTTTGCCAGAGGTTGCAACATGTTGAGAGAACCCGGTCAATAAAGTGTACTACCTCTTACCCCT

ABCA6 GENBANK: U66680

 TTTTCCCACAGGCTGCAGGGCAGGAAAGGTATTCCTCTTTGTTAACCTATAAGCTGCCCC GTGGCAGACGTTTACCCTCTATCACAGACCTTTCACAAATTAGAAGCAGTGAAAGCATAA CTTTAACCTGGAAGAATACAGCCTTTCTCCAGTGCACACTGGANAAGGTNTCCTTANAAC CTTCCTAAANAACAGGAAGTTAGGAAATTTTGAATGAAAANNNACCNCCCCCCCTCATTC AGGTGGAACCTTAAAACCTCAAACCTAGTAATTTTTTGTTGATCTCCTATAAAACTTATG TTTTATGTAATAATTAATAGTATGTTTAATTTTAAAGATCATTTAAAATTAACATCAGGT ATATTTTGTAAATTTAGTTAACAAATACATAAATTTTAAAATTATTCTTCCTCTCAAACA TAGGGGTGA TAGCAAACCTGTGATAAAGGCAATACAAAATATTAGTAAAGTCACCCAAAG AGTCAGGCACTGGGTATTGTGGAAATAAAACTATATAAACTTAA

ABCC4 GENBANK: U66682

ATGGATAAGTTTATACTAGTGTTGGCACATGGCGGCATGTATAGATATACTAGGAGGACC TAGTTGTATTCCTTGTATGAAAAAGCGTCCCTGGTACTACAATAAGTCTTTCGTGAAAGG AGTGTAATCCTAACAACAACTCAGGAAAGTATTTTGAAAAGAATACTGGATAAGGAAAAA GCTGTGGCCGTGATTCCTTGGATCGCAATACCCTTGGTTCCCCTTGGAATCATTTCATT TTTCTTCGGCGATATTTTTTGGAAACGTCAAGAGATGTGAAGCGCCTGGAATCTACAAGT CAGTATGGAAACTCGGGTTGGTATAGACATGCTAGCTAGTTTCCATTTATGCCATAAATT ACAGAGACCCCCTGAAATTCGGCAGACTCTGTCTTCCAGAATTTCTCTAACATTAGGTAA TTGAACGTATTGGCCATTATGAATCATTGTGTCCCTTAGAGCATGTGGAATTGATAGCCT GCAACGTGTAACTTTGCATTTGGAATAAGGAAGGAGTGAAGGCCATATGGGGAGTAATAT TCTACAGGAATGTCAGCACTGTGAAGACAGGGACTC

ABCA1 Acc.Nr.: AJ012376 GENBANK: HSA012376

CAAACATGTCAGCTGTTACTGGAAGTGGCCTGGCCTCTATTTATCTTCCTGATCCTGATC TCTGTTCGGCTGAGCTACCCACCCTATGAACAACATGAATGCCATTTTCCAAATAAAGCC ATGCCCTCTGCAGGAACACTTCCTTGGGTTCAGGGGATTATCTGTAATGCCAACAACCCC TGTTTCCGTTACCCGACTCCTGGGGAGGCTCCCGGAGTTGTTGGAAACTTTAACAAATCC ATTGTGGCTCGCCTGTTCTCAGATGCTCGGAGGCTTCTTTTATACAGCCAGAAAGACACC - AGCATGAAGGACATGCGCAAAGTTCTGAGAACATTACAGCAGATCAAGAAATCCAGCTCA AACTTGAAGCTTCAAGATTTCCTGGTGGACAATGAAACCTTCTCTGGGTTCCTGTATCACAACCTCTCTCCCAAAGTCTACTGTGGACAAGATGCTGAGGGCTGATGTCATTCTCCAC AAGGTATTTTTGCAAGGCTACCAGTTACATTTGACAAGTCTGTGCAATGGATCAAAATCA GAAGAGATGATTCAACTTGGTGACCAAGAAGTTTCTGAGCTTTGTGGCCTACCAAGGGAG AAACTGGCTGCAGCAGAGCGAGTACTTCGTTCCAACATGGACATCCTGAAGCCAATCCTG AGAACACTAAACTCTACATCTCCCTTCCCGAGCAAGGAGCTGGCCGAAGCCACAAAAACA ${\tt TTGCTGCATAGTCTTGGGACTCTGGCCCAGGAGCTGTTCAGCATGAGAAGCTGGAGTGAC}$ ATGCGACAGGAGGTGATGTTTCTGACCAATGTGAACAGCTCCAGCTCCTCCACCCAAATC TCTCTCAACTGGTATGAGGACAACAACTACAAAGCCCTCTTTGGAGGCAATGGCACTGAG GAAGATGCTGAAACCTTCTATGACAACTCTACAACTCCTTACTGCAATGATTTGATGAAG AATTTGGAGTCTAGTCCTCTTTCCCGCATTATCTGGAAAGCTCTGAAGCCGCTGCTCGTT GGGAAGATCCTGTATACACCTGACACTCCAGCCACAAGGCAGGTCATGGCTGAGGTGAAC CCCAAGATCTGGACCTTCATGGAGAACAGCCAAGAAATGGACCTTGTCCGGATGCTGTTG GACAGCAGGGACAATGACCACTTTTGGGAACAGCAGTTGGATGGCTTAGATTGGACAGCC CAAGACATCGTGGCGTTTTTGGCCAAGCACCCAGAGGATGTCCAGTCCAGTAATGGTTCT GTGTACACCTGGAGAGAGCTTTCAACGAGACTAACCAGGCAATCCGGACCATATCTCGC TTCATGGAGTGTGAACCTGAACAAGCTAGAACCCATAGCAACAGAAGTCTGGCTCATC *AACAAGTCCATGGAGCTGCTGGATGAGAGGAAGTTCTGGGCTGGTATTGTGTTCACTGGA* ATTACTCCAGGCAGCATTGAGCTGCCCCATCATGTCAAGTACAAGATCCGAATGGACATT GACAATGTGGAGAGGACAAATAAAATCAAGGATGGGTACTGGGACCCTGGTCCTCGAGCT GACCCCTTTGAGGACATGCGGTACGTCTGGGGGGGGCTTCGCCTACTTGCAGGATGTGGTG GAGCAGGCAATCATCAGGGTGCTGACGGGCACCGAGAAGAAACTGGTGTCTATATGCAA CAGATGCCCTATCCCTGTTACGTTGATGACATCTTTCTGCGGGTGATGAGCCGGTCAATGCCCCTCTTCATGACGCTGGCCTGGATTTACTCAGTGGCTGTGATCATCAAGGGCATCGTG TATGAGAAGGAGGCACGGCTGAAAGAGACCATGCGGATCATGGGCCTGGACAACAGCATC $\tt CTCTGGTTTAGCTGGTTCATTAGTAGCCTCATTCCTCTTGTGAGCGCTGGCCTGCTA$ GTGGTCATCCTGAAGTTAGGAAACCTGCTGCCCTACAGTGATCCCAGCGTGGTGTTTTGTC TTCCTGTCCGTGTTTGCTGTGGTGACAATCCTGCAGTGCTTCCTGATTAGCACACTCTTC TCCAGAGCCAACCTGGCAGCAGCCTGTGGGGGGCATCATCTACTTCACGCTGTACCTGCCC TACGTCCTGTGTGTGGCATGGCAGGACTACGTGGGCTTCACACTCAAGATCTTCGCTAGC CTGCTGTCTCCTGTGGCTTTTGGGTTTGGCTGTGAGTACTTTGCCCTTTTTGAGGAGCAG GGCATTGGAGTGCAGTGGGACAACCTGTTTGAGAGTCCTGTGGAGGAAGATGGCTTCAAT CTCACCACTTCGGTCTCCATGATGCTGTTTGACACCTTCCTCTATGGGGTGATGACCTGG TACATTGAGGCTGTCTTTCCAGGCCAGTACGGAATTCCCAGGCCCTGGTATTTTCCTTGC ACCAAGTCCTACTGGTTTGGCGAGGAAAGTGATGAGAAGAGCCACCCTGGTTCCAACCAG ATTCAGAACCTGGTAAAAGTCTACCGAGATGGGGTGAAGGTGGCTGTCGATGGCCTGGCA $\tt CTGAATTTTATGAGGGCCAGATCACCTCCTTCCTGGGCCACAATGGAGCGGGAAGACG$ ACCACCATGTCAATCCTGACCGGGTTGTTCCCCCCGACCTCGGGCACCGCCTACATCCTG GGAAAAGACATTCGCTCTGAGATGAGCACCATCCGGCAGAACCTGGGGGTCTGTCCCCAG CATAACGTGCTGTTTGACATGCTGACTGTCGAAGAACACATCTGGTTCTATGCCCGCTTG AAAGGGCTCTCTGAGAAGCACGTGAAGGCGGAGATGGAGCAGATGGCCCTGGATGTTGGT TTGCCATCAAGCAAGCTGAAAAGCAAAACAAGCCAGCTGTCAGGTGGAATGCAGAAAAG CTATCTGTGGCCTTGGCCTTTGTCGGGGGATCTAAGGTTGTCATTCTGGATGAACCCACA ${\tt GCTGGTGTGGACCCTTACTCCCGCAGGGGAATATGGGAGCTGCTGCTGAAATACCGACAA}$ GGCCGCACCATTATTCTCTCTACACACCACATGGATGAAGCGGACGTCCTGGGGGACAGG ATTGCCATCATCTCCCATGGGAAGCTGTGCTGTGTGGGGCTCCTCCCTGTTTCTGAAGAAC

 ${\tt TCCTGCAGAAACAGTAGTAGCACTGTGTCATACCTGAAAAAGGAGGACAGTGTTTCTCAG$ AGCAGTTCTGATGCTGGCCTGGGCAGCGACCATGAGAGTGACACGCTGACCATCGATGTC TCTGCTATCTCCAACCTCATCAGGAAGCATGTGTCTGAAGCCCGGCTGGTGGAAGACATA CTCTTTCATGAGATTGATGACCGGCTCTCAGACCTGGGCATTTCTAGTTATGGCATCTCA GAGACGACCCTGGAAGAAATATTCCTCAAGGTGGCCGAAGAGAGTGGGGTGGATGCTGAG ACCTCAGATGGTACCTTGCCAGCAAGACGAAACAGGCGGGCCTTCGGGGACAAGCAGAGC TGTCTTCGCCCGTTCACTGAAGATGATGCTGCTGATCCAAATGATCTGACATAGACCCA GAATCCAGAGAGACAGACTTGCTCAGTGGGATGGATGGCAAAGGGTCCTACCAGGTGAAA GGCTGGAAACTTACACAGCAACAGTTTGTGGCCCCTTTTGTGGAAGAGACTGCTAATTGCC AGACGGAGTCGGAAAGGATTTTTTGCTCAGATTGTCTTGCCAGCTGTGTTTGTCTGCATT GCCCTTGTGTTCAGCCTGATCGTGCCACCCTTTGGCAAGTACCCCAGCCTGGAACTTCAG CCCTGGATGTACAACGAACAGTACACATTTGTCAGCAATGATGCTCCTGAGGACACGGGA ACCCTGGAACTCTTAAACGCCCTCACCAAAGACCCTGGCTTCGGGACCCGCTGTATGGAA GGAAACCCAATCCCAGACACGCCCTGCCAGGCAGGGAGGAAGAGTGGACCACTGCCCCA GTTCCCCAGACCATCATGGACCTCTTCCAGAATGGGAACTGGACAATGCAGAACCCTTCA CCTGCATGCCAGTGTAGCAGCGACAAAATCAAGAAGATGCTGCCTGTGTGTCCCCCAGGG GCAGGGGGGCTGCCTCCACAAAGAAAACAAAACACTGCAGATATCCTTCAGGACCTG ACAGGAAGAACATTTCGGATTATCTGGTGAAGACGTATGTGCAGATCATAGCCAAAAGC TTAAAGAACAAGATCTGGGTGAATGAGTTTAGGTATGGCGGCTTTTCCCTGGGTGTCAGTCACCTAAAGCTGGCCAAGGACAGTTCTGCAGATCGATTTCTCAACAGCTTGGGAAGATTT ATGACAGGACTGGACACCAGAAATAATGTCAAGGTGTGGTTCAATAACAAGGGCTGGCAT GCAATCAGCTCTTTCCTGAATGTCATCAACAATGCCATTCTCCGGGCCAACCTGCAAAAG GGAGAGACCCTAGCCATTATGGAATTACTGCTTTCAATCATCCCCTGAATCTCACCAAG CAGCAGCTCTCAGAGGTGGCTCCGATGACCACATCAGTGGATGTCCTTGTGTCCATCTGT GTCATCTTTGCAATGTCCTTCGTCCCAGCCAGCTTTGTCGTATTCCTGATCCAGGAGCGGGTCAGCAAAGCAAAACACCTGCAGTTCATCAGTGGAGTGAAGCCTGTCATCTACTGGCTC TCTAATTTTGTCTGGGATATGTGCAATTACGTTGTCCCTGCCACACTGGTCATTATCATC TTCATCTGCTTCCAGCAGAAGTCCTATGTGTCCTCCACCAATCTGCCTGTGCTAGCCCTT CTACTTTTGCTGTATGGGTGGTCAATCACACCTCTCATGTACCCAGCCTCCTTTGTGTTC AAGATCCCCAGCACAGCCTATGTGGTGCTCACCAGCGTGAACCTCTTCATTGGCATTAAT GGCAGCGTGGCCACCTTTGTGCTGGAGCTGTTCACCGACAATAAGCTGAATAATATCAAT GATATCCTGAAGTCCGTGTTCTTGATCTTCCCACATTTTTGCCTGGGACGAGGGCTCATC GACATGGTGAAAAACCAGGCAATGGCTGATGCCCTGGAAAGGTTTGGGGAGAATCGCTTT GTGTCACCATTATCTTGGGACTTGGTGGGACGAAACCTCTTCGCCATGGCCGTGGAAGGG GTGGTGTTCTTCCTCATTACTGTTCTGATCCAGTACAGATTCTTCATCAGGCCCAGACCT GTAAATGCAAAGCTATCTCCTCTGAATGATGAAGATGAAGATGTGAGGCGGGAAAGACAG

AGAATTCTTGATGGTGGAGGCCAGAATGACATCTTAGAAATCAAGGAGTTGACGAAGATA TATAGAAGGAAGCGGAAGCCTGCTGTTGACAGGATTTGCGTGGGCATTCCTCCTGGTGAG TGCTTTGGGCTCCTGGGAGTTAATGGGGCTGGAAAATCATCAACTTTCAAGATGTTAACA GGAGATACCACTGTTACCAGAGGAGGATGCTTTCCTTAACAGAAATAGTATCTTATCAAACATCCATGAAGTACATCAGAACATGGGCTACTGCCCTCAGTTTGATGCCATCACAGAGCTG GTTGGCAAGGTTGGTGAGTGGGCGATTCGGAAACTGGGCCTCGTGAAGTATGGAGAAAAA TATGCTGGTAACTATAGTGGAGGCAACAAACGCAAGCTCTCTACAGCCATGGCTTTGATC GGCGGGCCTCCTGTGGTGTTTCTGGATGAACCCACCACAGGCATGGATCCCAAAGCCCGG CGGTTCTTGTGGAATTGTGCCCTAAGTGTTGTCAAGGAGGGGAGATCAGTAGTGCTTACATCTCATAGTATGGAAGAATGTGAAGCTCTTTGCACTAGGATGGCAATCATGGTCAATGGA AGGTTCAGGTGCCTTGGCAGTGTCCAGCATCTAAAAAATAGGTTTGGAGATGGTTATACA ATAGTTGTACGAATAGCAGGGTCCAACCCGGACCTGAAGCCTGTCCAGGATTTCTTTGGA CTTGCATTTCCTGGAAGTGTTCCAAAAGAGAAACACCGGAACATGCTACAATACCAGCTT CCATCTTCATTATCTTCTCTGGCCAGGATATTCAGCATCCTCTCCCAGAGCAAAAAGCGA CTCCACATAGAAGACTACTCTGTTTCTCAGACAACACTTGACCAAGTATTTGTGAACTTT GCCAAGGACCAAAGTGATGATGACCACTTAAAAGACCTCTCATTACACAAAAACCAGACA GTATGAAGAATCCTGTTCATACGGGGTGGCTGAAAGTAAAGAGGGACTAGACTTTCCTTT ${\it GCACCATGTGAAGTGTTGTGGAGAAAAGAGCCAGAAGTTGATGTGGGAAGAAGTAAACTG}$ GATACTGTACTGATACTATTCAATGCAATGCAATTCAATG

ABCD2 Acc.Nr.: AJ000327 GENBANK: HSALDR

AAAACACAACAGTGGAAGAGAAACGCTGCATACTATGGGACGCTGTAGGACTTTCTAAAA ${\it CATTTGCTGGGGATTTCTGTGAAGCATGATCTTTTAAACGAATTCTTTTGGAAGCCGGTT}$ TGGGTAACTGGGAAAATGACACATATGCTAAATGCAGCAGCTGATCGAGTGAAATGGACCAGATCGAGTGCTGCTAAGAGGGCTGCCTGCCTGGTGGCTGCGGCATATGCTCTGAAAACCCTCTATCCCATCATTGGCAAGCGTTTAAAGCAATCTGGCCACGGGAAGAAAAAAGCAGCAGCTTACCCTGCTGCAGAGAACACAGAAATACTGCATTGCACCGAGACCATTTGTGAAAAA CCTTCGCCTGGAGTGAATGCAGATTTCTTCAAACAGCTACTAGAACTTCGGAAAATTTTG GTGGAAAAGAAGCCTCGGACTTTCATCATCAAATTAATCAAGTGGCTTATGATTGCCATC CCTGCTACCTTCGTCAACAGTGCAATAAGGTACCTGGAATGCAAATTGGCTTTGGCCTTC AGAACTCGCCTAGTAGACCACGCCTATGAAACCTATTTTACAAATCAGACTTATTATAAA GTGATCAATATGGATGGGAGGCTGGCAAACCCTGACCAATCTCTTACGGAGGATATTATG ATGTTCTCCCAATCTGTGGCTCACTTGTATTCCAATCTGACCAAACCTATTTTAGATGTA ATGCTGACCTCCTATACACTCATTCAAACTGCTACATCCAGAGGAGCAAGCCCAATTGGG CCCACCCTACTAGCAGGACTTGTGGTGTATGCCACTGCTAAAGTGTTAAAAGCCTGTTCT CCCAAATTTGGCAAACTGGTGGCAGAGGAAGCACATAGAAAAGGCTATTTGCGGTATGTG

CACTCGAGAATTATAGCCAATGTAGAAGAAATTGCCTTTTACAGAGGACATAAGGTAGAA ATGAAACAACTTCAGAAAAGTTACAAAGCTTTAGCAGATCAGATGAACCTCATTTTATCC AAACGTTTGTGGTACATCATGATAGAACAGTTCCTGATGAAGTATGTTTGGAGCAGCAGT GGACTAATTATGGTGGCTATACCTATTATCACTGCAACTGGCTTTGCAGATGGTGAGGAT GGCCAAAAGCAAGTTATGGTTAGTGAACGGACAGAAGCCTTTACCACTGCTCGAAATTTA CTGGCCTCTGGAGCTGATGCTATTGAAAGGATTATGTCTTCATACAAAGAGGTCACTGAATTAGCAGGCTACACTGCTCGAGTGTACAATATGTTTTGGGTCTTTGATGAAGTAAAAAGA GGCATTTATAAGAGAACTGCTGTCATTCAAGAATCTGAAAGCCATAGCAAGAATGGAGCT AAGGTAGAATTACCTCTCAGTGACACATTGGCAATTAAAGGAAAAGTTATTGATGTGGATCACGGAATTATTTGTGAAAATGTTCCCATAATTACACCAGCAGGAGAAGTGGTGGCTTCCAGGCTAAACTTCAAAGTAGAAGAAGGAATGCATCTTTTGATAACTGGTCCCAATGGTTGT GGGAAAAGTTCTCTTCAGAATTCTAAGTGGGCTCTGGCCTGTGTATGAAGGAGTCCTCTATAAACCACCTCCTCAACATATGTTTTATATTCCACAAAGGCCATATATGTCTCTTGGA AGTCTTCGGGATCAAGTCATTTACCCTGATTCAGTGGATGATATGCATGATAAAGGTTAT ACAGACCAAGATCTGGAACGTATCCTACACAATGTCCATCTCTATCACATAGTTCAAAGA GAAGGAGGATGGGATGCTGTTATGGACTGGAAAGATGTCCTGTCAGGAGGGGAAAAGCAA AGAATGGGCATGGCTCGTATGTTTTATCATAAACCAAAATATGCCTTGCTGGATGAATGT ACCAGTGCTGTCAGCATTGATGTCGAAGGAAAGATATTTCAGGCTGCAAAAGGGGCTGGA ATTTCCTTACTGTCTATAACACACAGACCTTCTCTTTGGAAATACCACACACTTTATTA CAGTTTGATGGTGAAGGAGGTTGGCGCTTTGAACAATTGGATACTGCTATCCGTTTGACATTGAGTGAAGAAAACAAAAGCTAGAATCTCAGCTAGCTGGAATTCCCAAAATGCAGCAG AGACTCAATGAACTATGTAAAATTTTGGGAGAAGACTCAGTGCTGAAAACAATTAAAAAT GAAGATGAGACATCTTAATTTGTTTTGACATATTTTAAAAAGTTAATTATTAGATAAAGG $\tt CTCAAAGACATTCTGTTATACTGCATGAAGTATGTTAAGCTAAGCACAGAGAAAAAAAGG$ CAGCAAGACATGTTTTATAAGATTTTAGCATTAAGGAAGTATATGATCTGACTTTTCAGAAGAAAATAAACAAATGCATTATGTAAGGTCAGTCATTATGACTTATACTAATTCCTAGTG AAGGCCTAATGCACTTGTAAAACAGGATTTTCTAGGTGAATTCCTGATGAATACCAGATTTACTATGTATATGTGGTGTGTCTGAAGTTCTTAACAACATGGGCAATATTCTGGAAATG AAACAAGTTATAACTGAGCACCATTTGGGTTGATACCAAGTGCATAAGATTCAAACTTTG AGTGACATTTAGTCCATTTATGGTTGATATTAGGTTTAATACCTAGAATTCAAATTGATT ATTGCTAGTGGCCAACTAAACCTGTACAAAATAGCTGACAGTTTTATAACTAATTTCAAT ATAAAAATTGTTTTAATGGCATTTGTTGAAAGAAAAAGCATGGCTAAAATGTATCAAAT TAGTACAATCTTAAATATTTTTAATAAATCCTTTCATTTTAAAAAGAGAATTGCCAATAC AGAAAAGGAGTATCCAAACAATGTCTCAACCTGATAATTTCCTTAGCAGAATTACCTATT GCAACTTCTGTTCAGAAATACACAGCTTGTTTTTTTGCCCAAGGATGAGTCTACATTTTAGGAATAGTACTTTATAATTTACAATCCCCATTTACATCATCATTTCACCTTAATGTTGAGGAC AATGTTTTGAAACAAATACTATTTTTCCTACTTTTGCTTTTGAGAAAATTGACACTCAGAC

ABCB1 Acc.Nr. M14758 GENBANK: HUMMDR1

CCTACTCTATTCAGATATTCTCCAGATTCCTAAAGATTAGAGATCATTTCTCATTCTCCT AGGAGTACTCACTTCAGGAAGCAACCAGATAAAAGAGAGGTGCAACGGAAGCCAGAACAT TCCTCCTGGAAATTCAACCTGTTTCGCAGTTTCTCGAGGAATCAGCATTCAGTCAATCCG GGCCGGGAGCAGTCATCTGTGGTGAGGCTGATTGGCTGGGCAGGAACAGCGCCGGGGGCGTGGGCTGAGCACAGCGCTTCGCTCTCTTTGCCACAGGAAGCCTGAGCTCATTCGAGTAGCG GCTCTTCCAAGCTCAAAGAAGCAGAGGCCGCTGTTCGTTTCCTTTAGGTCTTTCCACTAA AGTCGGAGTATCTTCTTCCAAGATTTCACGTCTTGGTGGCCGTTCCAAGGAGCGCGAGGT CGGGATGGATCTTGAAGGGGACCGCAATGGAGGAGCAAAGAAGAAGAACTTTTTTAAACT ${\it GAACAATAAAAGTGAAAAAGATAAGAAGGAAAAGAAACCAACTGTCAGTGTATTTTCAAT$ GTTTCGCTATTCAAATTGGCTTGACAAGTTGTATATGGTGGTGGGAACTTTGGCTGCCAT CATCCATGGGGCTGGACTTCCTCTCATGATGCTGGTGTTTTGGAGAAATGACAGATATCTTTGCAAATGCAGGAAATTTAGAAGATCTGATGTCAAACATCACTAATAGAAGTGATATCAA ${\tt TGATACAGGGTTCTTCATGAATCTGGAGGAAGACATGACCAGGTATGCCTATTATTACAG$ TGGAATTGGTGCTGGGTGCTGGTTGCTGCTTACATTCAGGTTTCATTTTGGTGCCTGGC AGCTGGAAGACAAATACACAAAATTAGAAAACAGTTTTTCATGCTATAATGCGACAGGA GATAGGCTGGTTTGATGTGCACGATGTTGGGGGAGCTTAACACCCGACTTACAGATGATGTCTCTAAGATTAATGAAGTTATTGGTGACAAAATTGGAATGTTCTTTCAGTCAATGGCAACATTTTTCACTGGGTTTATAGTAGGATTTACACGTGGTTGGAAGCTAACCCTTGTGATTTTGGCCATCAGTCCTGTTCTTGGACTGTCAGCTGCTGTCTGGGCAAAGATACTATCTTCATTTACTGATAAAGAACTCTTAGCGTATGCAAAAGCTGGAGCAGTAGCTGAAGAGGTCTTGGCAGCAATTAGAACTGTGATTGCATTTGGAGGACAAAGAAGAACTTGAAAGGTACAACAA AAATTTAGAAGAAGCTAAAAGAATTGGGATAAAGAAAGCTATTACAGCCAATATTTCTAT AGGTGCTGCTTTCCTGCTGATCTATGCATCTTATGCTCTGGCCTTCTGGTATGGGACCACCTTGGTCCTCTCAGGGGAATATTCTATTGGACAAGTACTCACTGTATTCTTTTCTGTATT AATTGGGGCTTTTAGTGTTGGACAGGCATCTCCAAGCATTGAAGCATTTGCAAATGCAAG AGGAGCAGCTTATGAAATCTTCAAGATAATTGATAATAAGCCAAGTATTGACAGCTATTC GAAGAGTGGGCACAAACCAGATAATATTAAGGGAAATTTGGAATTCAGAAATGTTCACTT CAGTTACCCATCTCGAAAAGAAGTTAAGATCTTGAAGGCCTGAACCTGAAGGTGCAGAG TGGGCAGACGGTGGCCCTGGTTGGAAACAGTGGCTGTGGGAAGAGCACAACAGTCCAGCTGATGCAGAGGCTCTATGACCCCACAGAGGGGATGGTCAGTGTTGATGGACAGGATATTAG GACCATAAATGTAAGGTTTCTACGGGAAATCATTGGTGTGGTGAGTCAGGAACCTGTATT GTTTGCCACCACGATAGCTGAAAACATTCGCTATGGCCGTGAAAATGTCACCATGGATGA GATTGAGAAAGCTGTCAAGGAAGCCAATGCCTATGACTTTATCATGAAACTGCCTCATAA ATTTGACACCCTGGTTGGAGAGAGGGGGCCCAGTTGAGTGGTGGGCAGAAGCAGAGGAT

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ABCB4 Acc. Nr.: M23234 GENBANK: HUMMDR3

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ABCC2 Acc.Nr.: U49248 GENBANK: HSU49248

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CTAAGCAGGTATTCGTTGGTTTTCTTCTTATTCTAGCAGCCATAGAGCTGGCCCTTGTAC TCACAGAAGACTCTGGACAAGCCACAGTCCCTGCTGTTCGATATACCAATCCAAGCCTCT ACCTAGGCACATGGCTCCTGGTTTTGCTGATCCAATACAGCAGACAATGGTGTGTACAGA AAAACTCCTGGTTCCTGTCCCTATTCTGGATTCTCTCGATACTCTGTGGCACTTTCCAAT TTCAGACTCTGATCCGGACACTCTTACAGGGTGACAATTCTAATCTAGCCTACTCCTGCC TGTTCTTCATCTCCTACGGATTCCAGATCCTGATCTTTTCAGCATTTTCAGAAAATAATGAGTCATCAAATAATCCATCATCCATAGCTTCATTCCTGAGTAGCATTACCTACA GCTGGTATGACAGCATCATTCTGAAAGGCTACAAGCGTCCTCTGACACTCGAGGATGTCT GGGAAGTTGATGAAGAGATGAAAACCAAGACATTAGTGAGCAAGTTTGAAACGCACATGA AGAGAGAGCTGCAGAAAGCCAGGCGGCACTCCAGAGACGGCAGGAGAAGAGCTCCCAGC AGAACTCTGGAGCCAGGCTGCCTGGCTTGAACAAGAATCAGAGTCAAAGCCAAGATGCCC TTGTCCTGGAAGATGTTGAAAAGAAAAAAAGAAGTCTGGGACCAAAAAAGATGTTCCAA AATCCTGGTTGATGAAGGCTCTGTTCAAAACTTTCTACATGGTGCTCCTGAAATCATTCCTACTGAAGCTAGTGAATGACATCTTCACGTTTGTGAGTCCTCAGCTGCTGAAATTGCTGA TCTCCTTTGCAAGTGACCGTGACACATATTTGTGGATTGGATATCTCTGTGCAATCCTCT TATTCACTGCGGCTCTCATTCAGTCTTTCTGCCTTCAGTGTTATTTCCAACTGTGCTTCA AGCTGGGTGTAAAAGTACGGACAGCTATCATGGCTTCTGTATATAAGAAGGCATTGACCC TATCCAACTTGGCCAGGAAGGAGTACACCGTTGGAGAAACAGTGAACCTGATGTCTGTGG ATGCCCAGAAGCTCATGGATGTGACCAACTTCATGCACATGCTGTGGTCAAGTGTTCTAC AGATTGTCTTATCTTCTTCCTATGGAGAGAGTTGGGACCCTCAGTCTTAGCAGGTGTTGGGGTGATGGTGCTTGTAATCCCAATTAATGCGATACTGTCCACCAAGAGTAAGACCAACCTCCGGAAGAAGAGCTCAAGAACCTGCTGGCCTTTAGTCAACTACAGTGTGTAGTAA TATTCGTCTTCCAGTTAACTCCAGTCCTGGTATCTGTGGTCACATTTTCTGTTTATGTCCTGGTGGATAGCAACAATATTTTGGATGCACAAAAGGCCTTCACCTCCATTACCCTCTTCAATATCCTGCGCTTTCCCCTGAGCATGCTTCCCATGATGATCTCCTCCATGCTCCAGGCCA GTGTTTCCACAGAGCGGCTAGAGAAGTACTTGGGAGGGGGATGACTTGGACACATCTGCCA ${\tt TTCGACATGACTGCAATTTTGACAAAGCCATGCAGTTTTCTGAGGCCTCCTTTACCTGGG}$ AACATGATTCGGAAGCCACAGTCCGAGATGTGAACCTGGACATTATGGCAGGCCAACTTG $\tt TGGCTGTGATAGGCCCTGTCGGCTCTGGGAAATCCTCCTTGATATCAGCCATGCTGGGAG$ AAATGGAAAATGTCCACGGGCACATCACCATCAAGGGCACCACTGCCTATGTCCCACAGC AGTCCTGGATTCAGAATGGCACCATAAAGGACAACATCCTTTTTGGAACAGAGTTTAATG AAAAGAGGTACCAGCAAGTACTGGAGGCCTGTGCTCCTCCCAGACTTGGAAATGCTGC CTGGAGGAGATTTGGCTGAGATTGGAGGAGGGGTATAAATCTTAGTGGGGGTCAGAAGCCCCTGTCTGCAGTGGATGCTCATGTAGGAAAACATATTTTTAATAAGGTCTTGGGCCCCAATGGCCTGTTGAAAGGCAAGACTCGACTCTTGGTTACACATAGCATGCACTTTCTTCCTCAAGTGGATGAGATTGTAGTTCTGGGGAATGGAACAATTGTAGAGAAAGGATCCTACAGTG

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ABCD1 Acc.Nr.: Z21876 GENBANK: HSXLALDA

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ABCC1 Acc.Nr.: L05628 GENBANK: HUMMRPX

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ABCC5 GENBANK: AF104942

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TGGTGTTCTTCTGTGGGAATGATCGCAGGAGTCTTCCCGTGGTTCCTTGTGGCAGTGG GGCCCCTTGTCATCCTCTTTTCAGTCCTGCACATTGTCTCCAGGGTCCTGATTCGGGAGC TGAAGCGTCTGGACAATATCACGCAGTCACCTTTCCTCTCCCACATCACGTCCAGCATAC AGGGCCTTGCCACCATCCACGCCTACAATAAAGGGCAGGAGTTTCTGCACAGATACCAGG AGCTGCTGGATGACAACCAAGCTCCTTTTTTTTTTTTTACGTGTGCGATGCGGTGGCTGG CTGTGCGGCTGGACCTCATCAGCATCGCCCTCATCACCACCGGGGGCTGATGATCGTTCTTATGCACGGGCAGATTCCCCCAGCCTATGCGGGTCTCGCCATCTCTTATGCTGTCCAGT TAACGGGGCTGTTCCAGTTTACGGTCAGACTGGCATCTGAGACAGAAGCTCGATTCACCT CGGTGGAGGGATCAATCACTACATTAAGACTCTGTCCTTGGAAGCACCTGCCAGAATTA AGAACAAGGCTCCCTCCCCTGACTGGCCCCAGGAGGGAGAGGTGACCTTTGAGAACGCAG CTAAAGAGAAGATTGGGGATGGGGGGGACAGGATCAGGGAAGTCCTCGCTGGGGATGGCCCTCTTCCGTCTGGTGGAGTTATCTGGAGGCTGCATCAAGATTGATGGAGTGAGAATCA GTGATATTGGCCTTGCCGACCTCCGAAGCAACTCTCTATCATTCCTCAAGAGCCGGTGC TGTTCAGTGGCACTGTCAGATCAAATTTGGACCCCTTCAACCAGTACACTGAAGACCAGA ${\tt TTTGGGATGCCCTGGAGGGGCACACACATGAAAGAATGTATTGCTCAGCTACCTCTGAAAC}$ TTGAATCTGAAGTGATGGAGAATGGGGGATAACTTCTCAGTGGGGGAACGGCAGCTCTTGT GCATAGCTAGAGCCCTGCTCCGCCACTGTAAGATTCTGATTTTAGATGAAGCCACAGCTGCCATGGACACAGAGACAGACTTATTGATTCAAGAGACCATCCGAGAAGCATTTGCAGACTGTACCATGCTGACCATTGCCCATCGCCTGCACACGGTTCTAGGCTCCGATAGGATTATGG TGCTGGCCCAGGGACAGGTGGTGGAGTTTGACACCCCATCGGTCCTTCTGTCCAACGACA GTTCCCGATTCTATGCCATGTTTGCTGCTGCAGAGACAAGGTCGCTGTCAAGGGCTGAC CCCCTCATCGCGTCCTCCTACCGAAACCTTGCCTTTCTCGATTTTATCTTTCGCACAGCA ${\it GTTCCGGATTGGCTTGTGTGTTTCACTTTTAGGGAGGGCATATTTTGATTATTGTATTT}$ ATTCCATATTCATGTAAACAAAATTTAGTTTTTGTTCTTAATTGCACTCTAAAAGGTTCA GGGAACCGTTATTATAATTGTATCAGAGGCCTATAATGAAGCTTTATACGTGTAGCTATA TCTATATATATATCTGTACATAGCCTATATTTACAGTGAAAATGTAAGCTGTTTATTTTA TATTAAAATAAGCACTGTGCTAATAACAGTGCATATTCCTTTCTATCATTTTTGTACAGT TTGCTGTACTAGAGATCTGGTTTTGCTATTAGACTGTAGGAAGAGTAGCATTTCATTCTT CTCTAGCTGGTGGTTCACGGTGCCAGGTTTTCTGGGTGTCCAAAGGAAGACGTGTGGCA GAGACGGGTGGGCGGCTGGAGACCATGCAGAGCGCCGTGAGTTCTCAGGGCTCCTGCCTT $\tt TTTCACTCCCTCCATCAAGAATGGGGATCACAGAGACATTCCTCCGAGCCGGGGAGTTTC$ TTTCCTGCCTTCTTTTTTGCTGTTGTTTCTAAACAAGAATCAGTCTATCCACAGAGAGTCCCACTGCCTCAGGTTCCTATGGCTGGCCACTGCACAGAGCTCTCCAGGCTCCAAGACCT GTTGGTTCCAAGCCCTGGAGCCAACTGCTGCTTTTTGAGGTGGCACTTTTTCATTTGCCT ATTCCCACACCTCCACAGTTCAGTGGCAGGGCTCAGGATTTCGTGGGTCTGTTTTCCTTT

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ABCA5 Acc. Nr.: AF000148 GENBANK: HSAF000148 GCCAGAGGCGCTCTTAACGGCGTTTATGTCCTTTGCTGTCTGAGGGGCCTCAGCTCTGAC ${\it CAATCTGGTCTTCGTGGTCATTAGCATGGGCTTCGTGAGACAGATACAGCTTTTGCTC}$ TGGAAGACTGGACCCTGCGGAAAAGGCAAAAGATTCGCTTTGTGGTGGAACTCGTGTGG CCTTTATCTTTATTTCTGGTCTTGATCTGGTTAAGGAATGCCAACCGGTCTACAGCCATCATGAATGCCATTTCCCCAACAAGGCGATGCCCTCAGCAGGAATGCTGCCGTGGCTCCAG GGGATCTTCTGCAATGTGAACAATCCCTGTTTTCAAAGCCCCACCCCAGGAGAATCTCCT GGAATTGTGTCAAACTATAACAACTCCATCTTGGCAAGGGTATATCGAGATTTTCAAGAA CTCCTCATGAATGCACCAGAGAGCCAGCACCTTGGCCGTATTTGGACAGAGCTACACATCTTGTCCCAATTCATGGACACCCTCCGGACTCACCCGGAGAATTGCAGGAAGAGGAATA CGAATAAGGGATATCTTGAAAGATGAAGAAACACTGACACTATTTCTCATTAAAAACATC GCTCATGGAGTCCCGGACCTGGCGCTGAAGGACATCGCCTGCAGCGAGGCCCTCCTGGAG CGCTTCATCATCTTCAGCCAGAGACGCGGGGCAAAGACGGTGCGCTATGCCCTGTGCTCC CTCTCCCAGGGCACCCTACAGTGGATAGAAGACACTCTGTATGCCAACGTGGACTTCTTC AAGCTCTTCCGTGTGCTTCCCACACTCCTAGACAGCCGTTCTCAAGGTATCAATCTGAGA TCTTGGGGAGGAATATTATCTGATATGTCACCAAGAATTCAAGAGTTTATCCATCGGCCG AGTATGCAGGACTTGCTGTGGGTGACCAGGCCCCTCATGCAGAATGGTGGTCCAGAGACC TCTCGGGTGCTCTCCTTCAACTGGTATGAAGACAATAACTATAAGGCCTTTCTGGGGATT GACTCCACAAGGAAGGATCCTATCTATTCTTATGACAGAAGAACAACATCCTTTTGTAAT GCATTGATCCAGAGCCTGGAGTCAAATCCTTTAACCAAAATCGCTTGGAGGGCGGCAAAG CCTTTGCTGATGGGAAAAATCCTGTACACTCCTGATTCACCTGCAGCACGAAGGATACTG AAGAATGCCAACTCAACTTTTGAAGAACTGGAACACGTTAGGAAGTTGGTCAAAGCCTGG GAAGAAGTAGGGCCCCAGATCTGGTACTTCTTTGACAACAGCACACAGATGAACATGATC AGAGATACCCTGGGGAACCCAACAGTAAAAGACTTTTTGAATAGGCAGCTTGGTGAAGAA GGTATTACTGCTGAAGCCATCCTAAACTTCCTCTACAAGGGCCCTCGGGAAAGCCAGGCTGACGACATGGCCAACTTCGACTGGAGGGACATATTTAACATCACTGATCGCACCCTCCGC CTGGTCAATCAATACCTGGAGTGCTTGGTCCTGGATAAGTTTGAAAGCTACAATGATGAA ACTCAGCTCACCCAACGTGCCCTCTCTCTACTGGAGGAAAACATGTTCTGGGCCGGAGTG GTATTCCCTGACATGTATCCCTGGACCAGCTCTACCACCCCACGTGAAGTATAAGATC CGAATGGACATAGACGTGGTGGAGAAAACCAATAAGATTAAAGACAGGTATTGGGATTCT

GGTCCCAGAGCTGATCCCGTGGAAGATTTCCGGTACATCTGGGGCGGGTTTGCCTATCTG CAGGACATGGTTGAACAGGGGATCACAAGGAGCCAGGTGCAGGCGGAGGCTCCAGTTGGA ATCTACCTCCAGCAGATGCCCTACCCCTGCTTCGTGGACGATTCTTTCATGATCATCCTG AACCGCTGTTTCCCTATCTTCATGGTGCTGGCATGGATCTACTCTGTCTCCATGACTGTG AAGAGCATCGTCTTGGAGAAGGAGTTGCGACTGAAGGAGCCTTGAAAAATCAGGGTGTCTCCAATGCAGTGATTTGGTGTACCTGGTTCCTGGACAGCTTCTCCATCATGTCGATGAGC ATCTTCCTCCTGACGATATTCATCATGCATGTAAGAATCCTACATTACAGCGACCCATTC ATCCTCTTCCTGTTCTTGTTGGCTTTCTCCACTGCCACCATCATGCTGTGCTTTCTGCTC AGCACCTTCTTCTCCAAGGCCAGTCTGGCAGCCAGCCTGTAGTGGTGTCATCTATTTCACC CTCTACCTGCCACACCTCTGTGCTTCGCCTGGCAGGACCGCATGACCGCTGAGCTGAAGAAGGCTGTGAGCTTACTGTCTCCGGTGGCATTTGGATTTGGCACTGAGTACCTGGTTCGCTTTGAAGAGCAAGGCCTGGGGCTGCAGTGGAGCAACATCGGGAACAGTCCCACGGAAGGG ${\it GACGAATTCAGCTTCCTGCTGTCCATGCAGATGATGCTCCTTGATGCTGCTGTCTATGGC}$ TTACTCGCTTGGTACCTTGATCAGGTGTTTCCAGGAGACTATGGAACCCCACTTCCTTGG ${\tt TACTTTCTACAAGAGTCGTATTGGCTTGGCGGTGAAGGGTGTTCAACCAGAGAAGAA}$ AGAGCCCTGGAAAAGACCGAGCCCCTAACAGAGGAAACGGAGGATCCAGAGCACCCAGAA *AAGAATCTGGTAAAGATTTTTGAGCCCTCCGGCCGGCCAGCTGTGGACCGTCTGAACATC* ACCTTCTACGAGAACCAGATCACCGCATTCCTGGGCCACAATGGAGCTGGGAAAACCACC ACCTTGTCCATCCTGACGGGTCTGTTGCCACCAACCTCTGGGACTGTGCTCGTTGGGGGA AGGGACATTGAAACCAGCCTGGATGCAGTCCGGCAGAGCCTTGGCATGTGTCCACAGCAC AACATCCTGTTCCACCACCTCACGGTGGCTGAGCACATGCTGTTCTATGCCCAGCTGAAA GGAAAGTCCCAGGAGGAGGCCCAGCTGGAGATGGAAGCCATGTTGGAGGACACAGGCCTC CACCACAAGCGGAATGAAGAGGCTCAGGACCTATCAGGTGGCATGCAGAGAAAGCTGTCG GTTGCCATTGCCTTTGTGGGAGATGCCAAGGTGGTGATTCTGGACGAACCCACCTCTGGG GTGGACCCTTACTCGAGACGCTCAATCTGGGATCTGCTCCTGAAGTATCGCTCAGGCAGA $oldsymbol{ACCATCATCCACCACCATGGACGAGGCCGACCTCCTTGGGGACCGCATTGCC}$ ATCATTGCCCAGGGAAGGCTCTACTGCTCAGGCACCCCACTCTTCCTGAAGAACTGCTTT GGCACAGGCTTGTACTTAACCTTGGTGCGCAAGATGAAAAACATCCAGAGCCAAAGGAAA GGCAGTGAGGGGACCTGCAGCTGCTCGTCTAAGGGTTTCTCCACCACGTGTCCAGCCCAC GTTCTCCACCATGTTCCAGAGGCAAAGCTGGTGGAGTGCATTGGTCAAGAACTTATCTTC CTTCTTCCAAATAAGAACTTCAAGCACAGAGCATATGCCAGCCTTTTCAGAGAGCTGGAG GAGACGCTGGCTGACCTTGGTCTCAGCAGTTTTGGAATTTCTGACACTCCCCTGGAAGAG ATTTTTCTGAAGGTCACGGAGGATTCTGATTCAGGACCTCTGTTTGCGGGTGGCGCTCAG CAGAAAAGAGAAAACGTCAACCCCCGACACCCCTGCTTGGGTCCCAGAGAGAAGGCTGGA CAGACACCCCAGGACTCCAATGTCTGCTCCCCAGGGGCGCCGGCTGCTCACCCAGAGGGC CAGCCTCCCCAGAGCCAGAGTGCCCAGGCCCGCAGCTCAACACGGGGACACAGCTGGTC CTCCAGCATGTGCAGGCGCTGCTGGTCAAGAGATTCCAACACACCCATCCGCAGCCACAAG

GACTTCCTGGCGCAGATCGTGCTCCCGGCTACCTTTGTGTTTTTTGGCTCTGATGCTTTCT ATTGTTATCCCTCCTTTTGGCGAATACCCCGCTTTGACCCTTCACCCCTGGATATATGGG CAGCAGTACACCTTCTTCAGCATGGATGAACCAGGCAGTGAGCAGTTCACGGTACTTGCA GAGTACCCCTGTGGCAACTCAACACCCTGGAAGACTCCTTCTGTGTCCCCCAAACATCACC CAGCTGTTCCAGAAGCAGAAATGGACACAGGTCAACCCTTCACCATCCTGCAGGTGCAGC ACCAGGGAGAAGCTCACCATGCTGCCAGAGTGCCCGAGGGTGCCGGGGGCCTCCCGCCC CCCCAGAGAACACAGCGCAGCACGGAAATTCTACAAGACCTGACGGACAGGAACATCTCC GACTTCTTGGTAAAAACGTATCCTGCTCTTATAAGAAGCAGCTTAAAGAGCAAATTCTGG GTCAATGAACAGAGGTATGGAGGAATTTCCATTGGAGGAAAGCTCCCAGTCGTCCCCATC ACGGGGGAAGCACTTGTTGGGTTTTTAAGCGACCTTGGCCGGATCATGAATGTGAGCGGG GGCCCTATCACTAGAGAGGCCTCTAAAGAAATACCTGATTTCCTTAAACATCTAGAAACT GAAGACAACATTAAGGTGTGGTTTAATAACAAAGGCTGGCCATGCCCTGGTCAGCTTTCTC AATGTGGCCCACAACGCCATCTTACGGGCCAGCCTGCCTAAGGACAGGAGCCCCGAGGAG TATGGAATCACCGTCATTAGCCAACCCTGAACCTGACCAAGGAGCAGCTCTCAGAGATT ACAGTGCTGACCACTTCAGTGGATGCTGTGGTTGCCATCTGTGTGATTTTCTCCATGTCC $\tt TTCGTCCCAGCCAGCTTTGTCCTTTATTTGATCCAGGAGCGGGTGAACAAATCCAAGCAC$ CTCCAGTTTATCAGTGGAGTGAGCCCCACCACCTACTGGGTGACCAACTTCCTCTGGGACATCGTGAATTATTCCGTGAGTGCTGGGCTGGTGGGCATCTTCATCGGGTTTCAGAAG AAAGCCTACACTTCTCCAGAAAACCTTCCTGCCCTTGTGGGCACTGCTCCTGCTGTATGGA TGGGCGGTCATTCCCATGATGTACCCAGCATCCTTCCTGTTTGATGTCCCCAGCACAGCC TATGTGGCTTTATCTTGTGCTAATCTGTTCATCGGCATCAACAGCAGTGCTATTACCTTCATCTTGGAATTATTTGAGAATAACCGGACGCTGCTCAGGTTCAACGCCGTGCTGAGGAAG CTGCTCATTGTCTTCCCCCACTTCTGCCTGGGCCGGGGCCTCATTGACCTTGCACTGAGCCAGGCTGTGACAGATGTCTATGCCCGGTTTGGTGAGGAGCACTCTGCAAATCCGTTCCAC TGGGACCTGATTGGGAAGAACCTGTTTGCCATGGTGGTGGAAGGGGTGGTGCTTCCTCCTGACCCTGCTGGTCCAGCGCCACTTCTTCCTCTCCCAATGGATTGCCGAGCCCACTAAG GAGCCCATTGTTGATGAAGATGATGTGGCTGAAGAAAGACAAAGAATTATTACTGGT GGAAATAAAACTGACATCTTAAGGCTACATGAACTAACCAAGATTTATCCGGGCACCTCC AGCCCAGCAGTGGACAGGCTGTGTGTCGGAGTTCGCCCTGGAGAGTGCTTTGGCCTCCTG GGAGTGAATGGTGCCGGCAAAACAACCACATTCAAGATGCTCACTGGGGACAACACAGTG ACCTCAGGGGATGCCACCGTAGCAGGCAAGAGTATTTTAACCAATATTTCTGAAGTCCAT CAAAATATGGGCTACTGTCCTCAGTTTGATGCAATCGATGAGCTGCTCACAGGACGAGAA CATCTTTACCTTTATGCCCGGCTTCGAGGTGTACCAGCAGAAGAATCGAAAAGGTTGCA AGTGGGGGCAACAAGCGGAAACTCTCCACAGCCATCGCACTCATTGGCTGCCCACCGCTG GTGCTGCTGGATGAGCCCACCACAGGGATGGACCCCCAGGCACGCCGCATGCTGTGGAAC CAATGTGAGGCACTGTGTACCCGGCTGGCCATCATGGTAAAGGGCGCCTTTCGATGTATG

ABCG1 Acc.Nr.: U34919 GENBANK: HSU34919

GAATTCCGGGATGTGGAACGGTCGCAGGAGGCTGCTACAAGCCCCATGAGCAAGGCTGTT CCCACTGACAGAGCTTTCCCAGGATGACAGAGTGCGCTCTGCCTCTCTGGGGTGTGCT AGCCTACGAGGGGCAATCGTAAGGCGAATGTCACTGAAAGAACACAAGTGTCCTTAAACA TGGACTATCTGGGCTTTCTAGTGCTGAAATTCTTCCCACTCCCACTGCCCACTTCCCATT ATATAAAAACACAGTTGTTTCATGTTTTTGTTTCTTTACTGTTTTTCTTTGTT *AAGAATGCATTCATTTATTCAAAATTGTTTATTGTAGAATAATCAGGCATTGCGTGGATG* AGGTGGTGTCCAGCAACATGGAGGCCACTGAGACGGACCTGCTGAATGGACATCTGAAAA AAGTAGATAATAACCTCACGGAAGCCCAGCGCTTCTCCTCCTTGCCTCGGAGGGCAGCTG TGAACATTGAATTCAGGGACCTTTCCTATTCGGTTCCTGAAGGACCCTGGTGGAGGAAGA AAGGATACAAGACCCTCCTGAAAGGAATTTCCGGGAAGTTCAATAGTGGTGAGTTGGTGG GGGAGACGGGCATGAAGGGGGCCGTCCTCATCAACGGCCTGCCCCGGGACCTGCGCTGCT TCCGGAAGGTGTCCTGCTACATCATGCAGGATGACATGCTGCTGCCGCATCTCACTGTGC AGGAGGCCATGATGGTGTCGGCACATCTGAAGCTTCAGGAGAAGGATGAAGGCAGAAGGG AAATGGTCAAGGAGATACTGACAGCGCTGGGCTTGCTGTCTTGCGCCAACACGCGGACCG GGAGCCTGTCAGGTGGTCAGCGCAAGCGCCTGGCCATCGCGCTGGAGCTGGTGAACAACCCTCCAGTCATGTTCTTCGATGAGCCCACCAGCGCCTGGACAGCGCCTCCTGCTTCCAGG TGGTCTCGCTGATGAAAGGGCTCGCTCAAGGGGGTCGCTCCATCATTTGCACCATCCACCAGCCCAGCGCCAAACTCTTCGAGCTGTTCGACCAGCTTTACGTCCTGAGTCAAGGACAAT GTGTGTACCGGGGAAAAGTCTGCAATCTTGTGCCATATTTGAGGGATTTGGGTCTGAACTGCCCAACCTACCACAACCCAGCAGATTTTGTCATGGAGGTTGCATCCGGCGAGTACGGTG ATCAGAACAGTCGGCTGGTGAGAGCGGTTCGGGAGGGCATGTGTGACTCAGACCACAAGA GAGACCTCGGGGGTGATGCCGAGGTGAACCCTTTTCTTTGGCACCGGCCCTCTGAAGAGG TAAAGCAGACAAAACGATTAAAGGGGTTGAGAAAGGACTCCTCGTCCATGGAAGGCTGCC

ACAGCTTCTCTGCCAGCTGCCTCACGCAGTTCTGCATCCTCTTCAAGAGGACCTTCCTCA GCATCATGAGGGACTCGGTCCTGACACACCTGCGCATCACCTCGCACATTGGGATCGGCC TCCTCATTGGCCTGCTGTACTTGGGGATCGGGAACGAAGCCAAGAAGGTCTTGAGCAACT CCGGCTTCCTCTTCTCCCATGCTGTTCCTCATGTTCGCGGCCCTCATGCCTACTGTTCTGACATTTCCCCTGGAGATGGGAGTCTTTCTTCGGGAACACCTGAACTACTGGTACAGCC TGAAGGCCTACTACCTGGCCAAGACCATGGCAGACGTGCCCTTTCAGATCATGTTCCCAGTGGCCTACTGCAGCATCGTGTACTGGATGACGTCGCAGCCGTCCGACGCCGTGGCCTTTGTGCTGTTTGCCGCGCTGGGCACCATGACCTCCCTGGTGGCACAGTCCCTGGGCCTGCTGA ${\it TCGGAGCCGCCTCCACGTCCCTGCAGGTGGCCCACTTTCGTGGGCCCAGTGACAGCCATCC}$ CGGTGCTCCTGTTCTCGGGGTTCTTCGTCAGCTTCGACACCATCCCCACGTACCTACAGT GGATGTCCTACATCTCCTATGTCAGGTATGGGTTCGAAGGGGTCATCCTCTCCATCTATG GCTTAGACCGGGAAGATCTGCACTGTGACATCGACGAGGCGTGCCACTTCCAGAAGTCGG AGGCCATCCTGCGGGAGCTGGACGTGGAAAATGCCAAGCTGTACCTGGACTTCATCGTACTCGGGATTTTCTTCATCTCCCTCCGCCTCATTGCCTATTTTGTCCTCAGGTACAAATCC GGGCAGAGAGGTAAAACACCTGAATGCCAGGAAACAGGAAGATTAGACACTGTGGCCGAG ATCCAACCCCTAGAACCGCGTTGGGTTTGTGGGTGTCTCGTGCTCAGCCACTCTGCCCAG CTGGGTTGGATCTTCTCCATTCCCCTTTCTAGCTTAACTAGGAAGATGTAGGCAGATTGGTGGTTTTTTTTTTTTTTTAACATACAGAATTTTAAATACCACAACTGGGGCAGAATTTAAAGCTGCAACACACGCTGGTGATGAGAGGCTTCCTCAGTCCAGTCGCTCCTTAGCACCACACCGTGGGTCCTGGATGGGGAACTGCAAGCAGCCTCTCAGCTGATGGCTGCGCAGTC AGATGTCTGGTGGCAGAGAGTCCGAGCATGGAGCGATTCCATTTT

ABCA3 Acc.Nr: U78735 GENBANK: HSU78735

CCGCCCGGCGCCCAGGCTCGGTGCTGGAGAGTCATGCCTGTGAGCCCTGGGCACCTCCT GATGTCCTGCGAGGTCACGGTGTTCCCAAACCTCAGGGTTGCCCTGCCCCACTCCAGAGG CTCTCAGGCCCCACCCCGGAGCCCTCTGTGCGGAGCCGCCTCCTCCTGGCCAGTTCCCCAGTAGTCCTGAAGGGAGACCTGCTGTGTGGAGCCTCTTCTGGGACCCAGCCATGAGTGTGG AGCTGAGCAACTGAACCTGAAACTCTTCCACTGTGAGTCAAGGAGGCTTTTCCGCACATG AAGGACGCTGAGCGGGAAGGACTCCTCTCTGCCTGCAGTTGTAGCGAGTGGACCAGCACC CACGTCTGCACACCTCGCCCTCTTTACACTCAGTTTTCAGAGCACGTTTCTCCTATTTCCTGCGGGTTGCAGCGCCTACTTGAACTTACTCAGACCACCTACTTCTCTAGCAGCACTGGGCGTCCCTTTCAGCAAGACGATGGCTGTGCTCAGGCAGCTGGCGCTCCTCCTGGAAGAA $\tt CTACACCCTGCAGAAGCGGAAGGTCCTGGTGACGGTCCTGGAACTCTTCCTGCCATTGCT$ GTTTCCTGGGATCCTCATCTGGCTCCGCTTGAAGATTCAGTCGGAAAATGTGCCCAACGC CACCATCTACCCGGGCCAGTCCATCCAGGAGCTGCCTCTGTTCTTCACCTTCCCTCCGCCAGGAGACACCTGGGAGCTTGCCTACATCCCTTCTCACAGTGACGCTGCCAAGACCGTCAC TGAGACAGTGCGCAGGGCACTTGTGATCAACATGCGAGTGCGCGGCTTTCCCTCCGAGAA GGACTTTGAGGACTACATTAGGTACGACAACTGCTCGTCCAGCGTGCTGGCCGCCGTGGT

CTTCGAGCACCCCTTCAACCACAGCAGGAGCCCCTGCCGCTGGCGGTGAAATATCACCTACGGTTCAGTTACACACGGAGAAATTACATGTGGACCCAAACAGGCTCCTTTTTCCTGAA AGAGACAGAAGGCTGGCACACTACTTCCCTTTTCCCGCTTTTCCCAAACCCAGGACCAAG GGAACTAACATCCCCTGATGGCGGAGAACCTGGGTACATCCGGGAAGGCTTCCTGGCCGT GCAGCATGCTGTGGACCGGGCCATCATGGAGTACCATGCCGATGCCGCCACACGCCAGCT GTTCCAGAGACTGACGGTGACCATCAAGAGGTTCCCGTACCCGCCGTTCATCGCAGACCC CTTCCTCGTGGCCATCCAGTACCAGCTGCCCCCTGCTGCTGCTGCTCAGCTTCACCTACAC GCGCATGATGGGGCTCAGCAGCTGGCTGCACTGGAGTGCCTGGTTCCTCTTGTTCTTCCTCTTCCTCCTCATCGCCGCCTCCTTCATGACCCTGCTCTTCTGTGTCAAGGTGAAGCCAAACATCTCTACCATCTCCTTCAGCTTCATGGTCAGCACCTTCTTCAGCAAAGCCAACATGGC AGCAGCCTTCGGAGGCTTCCTCTACTTCTTCACCTACATCCCCTACTTCTTCGTGGCCCCTCGGTACAACTGGATGACTCTGAGCCAGAAGCTCTGCTCCTGCCTCCTGTCTAATGTCGC CATGGCAATGGGAGCCCAGCTCATTGGGAAATTTGAGGCGAAAGGCATGGGCATCCAGTG GCGAGACCTCCTGAGTCCCGTCAACGTGGACGACGACTTCTGCTTCGGGCAGGTGCTGGG GATGCTGCTGGACTCTGTGCTCTATGGCCTGGTGACCTGGTACATGGAGGCCGTCTT CCCAGGGCAGTTCGGCGTGCCTCAGCCCTGGTACTTCTTCATCATGCCCTCCTATTGGTGTGGGAAGCCAAGGGCGGTTGCAGGGAAGGAGGAAGAAGACAGTGACCCCGAGAAAGCACT CAGAAACGAGTACTTTGAAGCCGAGCCAGAGGACCTGGTGGCGGGGATCAAGATCAAGCA CCTGTCCAAGGTGTTCAGGGTGGGAAATAAGGACAGGGGGGGCCGTCAGAGACCTGAACCTCAACCTGTACGAGGGACAGATCACCGTCCTGCTGGGCCACAACGGTGCCGGGAAGACCAC CACCCTCTCCATGCTCACAGGTCTCTTTCCCCCCACCAGTGGACGGGCATACATCAGCGG GTATGAAATTTCCCAGGACATGGTTCAGATCCGGAAGAGCCTGGGCCTGTGCCCGCAGCA CGACATCCTGTTTGACAACTTGACAGTCGCAGAGCACCTTTATTTCTACGCCCAGCTGAAGGGCCTGTCACGTCAGAAGTGCCCTGAAGAAGTCAAGCAGATGCTGCACATCATCGGCCTGGAGGACAAGTGGAACTCACGGAGCCGCTTCCTGAGCGGGGGCATGAGGCGCAAGCTCTC CATCGGCATCGCCCTCATCGCAGGCTCCAAGGTGCTGATACTGGACGAGCCCACCTCGGGCATGGACGCCATCTCCAGGAGGGCCATCTGGGATCTTCTTCAGCGGCAGAAAAGTGACCG CACCATCGTGCTGACCACCCCACTTCATGGACGAGGCTGACCTGCTGGGAGACCGCATCGCCATCATGGCCAAGGGGGGGCTGCAGTGCTGCGGGTCCTCGCTGTTCCTCAAGCAGAAATA CGGTGCCGGCTATCACATGACGCTGGTGAAGGACCCGCACTGCAACCCGGAAGACATCTCCCAGCTGGTCCACCACCACGTGCCCAACGCCACGCTGGAGCAGCGCTGGGGCCGAGCTGTCTTCATCCTTCCCAGAGAGAGCACGCACAGGTTTGAAGGTCTCTTTGCTAAACTGGA AGTCTTCCTTCGGGTCGGGAAGCTGGTGGACAGCAGTATGGACATCCAGGCCATCCAGCT CCCTGCCTGCAGTACCAGCACGAGGGGGGGGGCGCCAGCGACTGGGCTGTGGACAGCAACCTTGTCAAGCTCAACACTGGGCTCGCCCTGCACTGCCAGCAATTCTGGGCCATGTTCCTGAA

GAAGGCCGCATACAGCTGGCGCGGGTGGAAAATGGTGGCGGCACAGGTCCTGGTGCCTCT GACCTGCGTCACCCTGGCCCTCCTGGCCATCAACTACTCCTCGGAGCTCTTCGACGACCC CATGCTGAGGCTGACCTTGGGCGAGTACGGCAGAACCGTCGTGCCCTTCTCAGTTCCCGG GACCTCCCAGCTGGGTCAGCAGCTGTCAGAGCCATCTGAAAGACGCACTGCAGGCTGAGGG ACAGGAGCCCCGCGAGGTGCTCGGTGACCTGGAGGAGTTCTTGATCTTCAGGGCTTCTGT GGAGGGGGGGCTTTAATGAGCGGTGCCTTGTGGCAGCGTCCTTCAGAGATGTGGGAGA GCGCACGGTCGTCAACGCCTTGTTCAACAACCAGGCGTACCACTCTCCAGCCACTGCCCT GGCCGTCGTGGACAACCTTCTGTTCAAGCTGCTGTGCGGGCCTCACGCCTCCATTGTGGTCTCCAACTTCCCCCAGCCCCGGAGCGCCCTGCAGGCTGCCAAGGACCAGTTTAACGAGGG CCGGAAGGGATTCGACATTGCCCTCAACCTGCTCTTCGCCATGGCATTCTTTGGCCAGCACGTTCTCCATCCTGGCGGTCAGCGAGAGGGCCGTGCAGGCCAAGCATGTGCAGTTTGTGAG GGACGGCCACATGGCTGACACCCTGCTGCTGCTCCTGCTCTACGGCTGGGCCATCATCCC ${\tt CCTCATGTACCTGATGAACTTCTTCTTCTTGGGGGGGGCCACTGCCTACACGAGGCTGAC}$ CATCTTCAACATCCTGTCAGGCATCGCCACCTTCCTGATGGTCACCATCATGCGCATCCC AGCTGTAAAACTGGAAGAACTTTCCAAAACCCTGGATCACGTGTTCCTGGTGCTGCCCAA CCACTGTCTGGGGATGGCAGTCAGCAGTTTCTACGAGAACTACGAGACGCGGAGGTACTG CACCTCCTCCGAGGTCGCCCCACTACTGCAAGAATATAACATCCAGTACCAGGAGAA CTTCTATGCCTGGAGCGCCCGGGGGGTCGGCCGGTTTGTGGCCTCCATGGCCGCCTCAGGGTGCGCCTACCTCATCCTGCTCTTCCTCATCGAGACCAACCTGCTTCAGAGACTCAGGGG CATCCTCTGCGCCCTCCGGAGGAGGCGGACACTGACAGAATTATACACCCGGATGCCTGT GCTTCCTGAGGACCAAGATGTAGCGGACGAGGACCCGCATCCTGGCCCCCAGCCCGGA CTCCCTGCTCCACACCCTCTGATTATCAAGGAGCTCTCCAAGGTGTACGAGCAGCGGGTGCCCCTCCTGGCCGTGGACAGGCTCTCCCTCGCGGTGCAGAAAGGGGAGTGCTTCGGCCT GCTGGGCTTCAATGGAGCCGGGAAGACCACGACTTTCAAAATGCTGACCGGGGAGGAGAG CCTCACTTCTGGGGATGCCTTTGTCGGGGGTCACAGATCAGCTCTGATGTCGGAAAGGT GCGGCAGCGGATCGGCTACTGCCCGCAGTTTGATGCCTTGCTGGACCACATGACAGGCCG GGAGATGCTGGTCATGTACGCTCGGCTCCGGGGCATCCCTGAGCGCCCACATCGGGGCCTG CGTGGAGAACACTCTGCGGGGCCTGCTGCTGGAGCCACATGCCAACAAGCTGGTCAGGAC GTACAGTGGTGAACAAGCGGAAGCTGAGCACCGGCATCGCCCTGATCGGAGAGCCTGCTGTCATCTTCCTGGACGAGCCGTCCACTGGCATGGACCCCGTGGCCCGGCGCCTGCTTTG GGACACCGTGGCACGAGCCCGAGAGTCTGGCAAGGCCATCATCACCTCCCACAGCAT GGAGGAGTGTGAGGCCTGTGCACCCGGCTGGCCATCATGGTGCAGGGGCAGTTCAAGTG CCTGGGCAGCCCCAGCACCTCAAGAGCAAGTTCGGCAGCGGCTACTCCCTGCGGGCCAA GGTGCAGAGTGAAGGGCAACAGGAGGCGCTGGAGGAGTTCAAGGCCTTCGTGGACCTGAC CTTTCCAGGCAGCGTCCTGGAAGATGAGCACCAAGGCATGGTCCATTACCACCTGCCGGGCCGTGACCTCAGCTGGGCGAAGGTTTTCGGTATTCTGGAGAAAGCCAAGGAAAAGTACGGCGTGGACGACTACTCCGTGAGCCAGATCTCGCTGGAACAGGTCTTCCTGAGCTTCGCCCA

Fragment 640918

- 1 GAGATCCTGAGGCTTTTCCCCCAGGCTGCTCAGCAGGAAAGGTTCTCCTCCCTGATGGTC
- 61 TATAAGTTGCCTGTTGAGGATGTGCGACCTTTATCACAGGCTTTCTTCAAATTAGAGATA
- 121 GTTAAACAGAGTTTCGACCTGGAGGAGTACAGCCTCTCACAGTCTACCCTGGAGCAGGTT
- 181 TTCCTGGAGCTCTCCAAGGAGCAGGAGCTGGGTGATCTTGAAGAGGACTTTGATCCCTCG
- 241 GTGAAGTGGAAACTCCTCCTGCAGGAAGAGCCTTAAAGCTCCAAATACCCTATATCTTTC
- 301 TTTAATCCTGTGACTCTTTTAAAGATAATATTTTATAGCCTTAATATGCCTTATATCAGA
- 361 GGTGGTACAAAATGCATTTGAAACTCATGCAATAATTATC

Fragment 698739

- 1 GCTCTCCACACAGAGATTTTGAAGCTTTTCCCACAGGCTGCTTGGCAGGAAAGATATTCC
- 61 TCTTTAATGGCGTATAAGTTACCTGTGGAGGATGTCCACCCTCTATCTCGGGCCTTTTTC
- 121 AAGTTAGAGGCGATGAAACAGACCTTCAACCTGGAGGAATACAGCCTCTCTCAGGCTACC
- 181 TTGGAGCAGGTATTCTTAGAACTCTGTAAAGAGCAGGAGCTGGGAAATGTTGATGATAAA
- 241 ATTGATACAACAGTTGAATGGAAACTTCTCCCACAGGAAGACCCTTAAAATGAAGAACCT
- $301\ CCTAACATTCAATTTTAGGTCCTACTACATTGTTAGTTTCCATAATTCTACAAGAATGTT$
- 361 TCCTTTTACTTCAGTTAACAAAGAAAACATTTAATAACATTCAATAATGATTACAGTT
 421 TTCATTTTTAAAAAATTTAGGATGAAGGAAACAAGGAAATATAGGGAAAAGTAGTAGACAA
- 481 AATTAACAAAATCAGACATGTTATTCATCCCCAACATGGGTCTATTTTGTGCTTAAAAAT
- 541 AATTTAAAAATCATACAATATTAGGTTGGTTATCG

Fragment 990006

- 1 GTGGAAGATGTGCAACCTTTAGCCCAAGCTTTCTTCAAATTAGAGAAGGTTAAACAGAGC
- 61 TTTGACCTAGAGGAGTACAGCCTCTCACAGTCTACCCTGGAGCAGGTTTTCCTGGAGCTC
- 121 TCCAAGGAGCAGGAGCTGGGTGATTTTGAGGAGGATTTTGATCCCTCAGTGAAGTGGAAG

- 181 CTCCTCCCCAGGAAGAGCCTTAAAACCCCAAATTCTGTGTTCCTGTTTAAACCCGTGGT
- 241 TTTTTTTAAATACATTTATTTTTTATAGCAGCAATGTTCTATTTTTAGAAACTATATTATA Fragment 1133530
- 1 TTTTCAGTTG CATGTAATAC CAAGAAATCG AATTGTTTTC CGGTTCTTAT
- 51 GGGAATTGTT AGCAATGCCC TTATTGGAAT TTTTAACTTC ACAGAGCTTA
- 101 TTCAAATGGA GAGCACCTTA TTTTTTCGTG ATGACATAGT GCTGGATCTT
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Huwhite2

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